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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT) (51) International Patent Classification 6: (11) International Publication Number: WO 99/04266 G01N 33/68 (43) International Publication Date: 28 January 1999 (28.01.99) PCT/GB98/02115 (21) International Application Number: (81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR, (22) International Filing Date: 16 July 1998 (16.07.98) (30) Priority Data: 9714971.0 16 July 1997 (16.07.97) GB

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BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, GM, HR, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

Published

Without international search report and to be republished upon receipt of that report.

(54) Title: ASSAYS, THERAPEUTIC METHODS AND MEANS

(57) Abstract

The interaction of ATM and related protein kinases such as ATR and DNA-PK with p53 is disclosed, in particular the phosphorylation of Ser15 and Thr18 by these proteins. The activity of the proteins is shown to increase in the presence of DNA. Assays for modulators of phosphorylation by the interaction between the proteins and p53 or other proteins having similar phosphorylation sites are provided. Methods of purifying ATM or ATR employing DNA or NTA are also disclosed.

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ASSAYS, THERAPEUTIC METHODS AND MEANS
The present invention relates to screening methods,
peptides, mimetics, and methods of use based on the
surprising discovery and characterisation of an
interaction between known proteins, and thus numerous
cellular processes of interest in therapeutic contexts.
The proteins in question are ATM and p53, and the
inventors have found that ATM phosphorylates p53 at a
number of specific sites. This interaction is observed
with other related proteins with associated kinase
activity, in particular ATR and DNA-PK, and other
proteins having similar phosphorylation sites to p53.
Further aspects of the present invention are founded on
the discovery that ATM binds DNA and that such binding
has an effect on phosphorylation of p53 by ATM.

Ataxia-telangiectasia (A-T) is a human autosomal recessive disorder characterised by a number of debilitating symptoms, including a progressive cerebellar degeneration, occulocutaneous telangiectasia, growth 20 retardation, immune deficiencies and certain characteristics of premature ageing (reviewed in Jackson, 1995; Meyn, 1995; Shiloh, 1995). A-T patients exhibit an approximately 100-fold increased incidence of cancer, with patients being particularly predisposed to 25 malignancies of lymphoid origin. Furthermore, A-T heterozygotes, which comprise ~1% of the population, are reported to exhibit a higher incidence of breast cancer (Easton, 1994; Meyn, 1995), although this remains controversial (Fitzgerald et al., 1997). At the cellular 30 level, A-T is characterised by a high degree of chromosomal instability, radioresistant DNA synthesis, and hypersensitivity to ionising radiation (IR) and radiomimetic drugs. In addition, A-T cells are defective in the radiation induced G1-S, S, and G2-M cell cycle 35 checkpoints that are thought to arrest the cell cycle in response to DNA damage in order to allow repair of the

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genome prior to DNA replication or mitosis (Halazonetis et al., 1993; Beamish et al., 1994; Beamish and Lavin, 1994; Khanna et al., 1995; Barlow et al., 1996; Xu and Baltimore, 1996). A-T cells exhibit deficient or severely delayed induction of p53 in response to IR (Kastan et al., 1992; Khanna and Lavin, 1993; Lu and Lane, 1993; Xu and Baltimore, 1996). p53 mediated transcriptional activation of p21/WAF1/CIP1 and Gadd45, and the subsequent inhibition of G1 cyclin-dependent kinases, are also defective in A-T cells following IR exposure (Artuso et al., 1995; Khanna et al., 1995). Lu and Lane, 1993, however, reported very little difference in the p53 response from normal and A-T cells.

- Furthermore, yeast have an ATM homologue (Mec1) but do not have p53 (Goffeau et al.). The best data for a possible substrate for Mec1p is Spk1/Rad53 (Sun et al; Sanchez et al.)
- The gene mutated in A-T patients, termed ATM (A-T 20 mutated), has been mapped and its cDNA cloned (Savitsky et al., 1995a; Savitsky et al., 1995b). analyses reveal that the ATM gene encodes a ~350 kDa polypeptide that is a member of the phosphatidylinositol (PI) 3-kinase family of proteins by virtue of a putative 25 kinase domain in its carboxyl-terminal region (Savitsky et al., 1995a; Savitsky et al., 1995b). Classical PI 3-kinases, such as PI 3-kinase itself, are involved in signal transduction and phosphorylate inositol lipids that act as intracellular second messengers (reviewed in 30 Kapeller and Cantley, 1994). ATM bears sequence similarity with a subset of the PI 3-kinase protein family that comprises proteins which, like ATM, are involved in cell cycle control and/or in the detection and signalling of DNA damage (for reviews see Hunter, 35 1995; Keith and Schreiber, 1995; Zakian, 1995; Jackson, Included in this sub-group are Saccharomyces

cerevisiae Torlp and Tor2p and their mammalian homologue FRAP, which control progression into S-phase and, at least in part, function by regulating translation (Brown and Schreiber, 1996). Also in this sub-group is the DNA dependent protein kinase (DNA-PK) catalytic subunit 5 (DNA-PKcs), defects in which lead to sensitivity to IR and an inability to perform site-specific V(D)J recombination (reviewed in Jackson and Jeggo, 1995; Jackson, 1996). Other members of the ATM sub-group of the PI 3-kinase family that have been identified include 10 S. cerevisiae Tellp and Meclp, together with the Meclp homologues of Schizosaccharomyces pombe (rad3), Drosophila melanogaster (mei-41) and humans FRP1/ATR; (Keith and Schreiber, 1995; Zakian, 1995; Jackson, 1996). As with ATM, defects in these proteins lead to genomic 15 instability, hypersensitivity towards DNA damaging agents and defects in DNA damage-induced cell cycle checkpoint controls.

- ATM is most similar to *S. cerevisiae* Tellp, which has not been shown to have any biochemical function so far (identity and similarity are 45% and 66%, respectively). ATM is much further diverged from DNA-PKcs (28% identical and 51% similar), with essentially the same homology to PI 3-kinase (a bona fide lipid kinase: 24% identical and 51% similar). Thus, from the sequence comparisons alone, one could not predict that ATM would be a protein kinase akin to DNA-PKcs or a lipid kinase akin to PI 3-kinase.
- Although genetic data indicate an involvement of ATM-like proteins in DNA damage recognition and its repair, the mechanisms by which these proteins function are not well understood. Much is known about the clinical symptoms and cellular phenotypes that arise from mutations in ATM, but little is known about the mechanisms by which the ATM protein functions. Recent studies have revealed that, like DNA-PKcs, ATM is expressed ubiquitously and is

localised predominantly in the cell nucleus (Chen and Lee, 1996; Lakin et al., 1996; Brown et al., 1997; Watters et al., 1997).

The realisation that ATM is a member of the PI 3-kinase 5 family has suggested to some that the primary function of ATM might phosphorylate inositol phospholipids. et al (1995 Science 268, 1749-1753), for example, do not discuss protein phosphorylation. Indeed, several lines of evidence suggest that ATM might have functioned in a 10 very different way from that which we have established For example, defective protein tyrosine phosphorylation and calcium mobilization in response to the triggering of B-cells and T-cells of A-T patients support the idea of defects in intra-cytoplasmic 15 signalling pathways in A-T cells (cited in the Savitski Science paper 1995). These data are provided in the paper Khanna et al (1997; J. Biol. Chem.). also summarises a variety of other data suggesting different ways in which ATM might function. 20

Savitsky et al (Science 1995) state that the insulin-dependent diabetes observed in some A-T patients could reflect ATM acting in an analogous way to PI 3-kinase affecting glucose transport by insulin. They also discuss PI 3-kinase in terms of controlling apoptosis as a paradigm for ATM, ie. one can explain many of the features of A-T by suggesting that it works analogously to PI 3-kinase.

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Some A-T cells have been shown to be complemented by a gene called ATDC, whose product interacts with an intermediate filament protein called vimentin, which is cytoplasmic (Brzoska et al; PNAS). They state that A-T cell lines have aberrantly aggregated actin filaments, suggesting the role of ATM lies in the cytoplasm.

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We have purified ATM. We report that, ATM binds to DNA and possesses an associated protein kinase activity that is stimulated by DNA. Furthermore, we show that ATM serves as a kinase for p53 and that the sites of phosphorylation reside in functionally important regions of the p53 polypeptide. These sites are Ser15 and Thr18. We also show that DNA-PK is also capable of phosphorylating the Ser15 and Thr18 sites of p53, and that ATR phosphorylates Ser15. Further, we show that phosphorylation of these sites of p53 disrupts the interaction of p53 with Mdm-2, a protein which targets p53 for degradation within the cell.

By targeting these sites, ATM may activate p53 for DNA binding and/or cause disassociation of Mdm-2, thus stabilising p53 (leading to increased amounts of the protein) and would allow it to activate transcription.

Thr18 of p53 has to our knowledge never been shown to be phosphorylated in vivo or in vitro. This site does not conform to a characterized DNA-PK consensus phosphorylation site. Thus, our finding of phosphorylation here is totally unexpected.

- Ser15 is phosphorylated by DNA-PK, but nonetheless its phosphorylation by ATM is also surprising, particularly since there are no data indicating its phosphorylation in reponse to DNA damage being altered in A-T cells.
- Based on this and other work described below, the present invention in various aspects provides for modulation of interaction between ATM (and ATR) and p53, particularly phosphorylation of p53 by ATM and ATR, and DNA binding by these proteins, which is further shown to have a potentiating effect on phosphorylation of p53.

Various aspects of the present invention provide for the

use of ATM (or related kinases such as ATR or DNA-PK) and p53, with or without DNA, in screening methods and assays for agents which modulate interaction between ATM and p53, particularly phosphorylation of p53 by ATM.

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Further aspects provide for modulation of interaction between ATM, or related kinases such as ATR or DNA-PK, and other molecules including a phosphorylation site homologous to those in p53 which are phosphorylated by ATM, and use of these molecules in screening methods and assays for useful agents. For simplicity, much of the present disclosure refers to ATM and p53. However, unless the context requires otherwise, every such reference should be taken to be equally applicable to the interaction between ATM and other molecules including a site homologous to one of those in p53 phosphorylated by Similarly, based on the disclosure herein, the invention extends to the use of other protein kinases which have an associated protein kinase activity capable of phosphorylating sites of p53, in particular Ser15 and Typically, the protein kinase domain of these other kinases will share at least 30% amino acid sequence identity with the corresponding domain of ATM, more preferably at least 35% sequence identity, more preferably at least 40% sequence identity, more preferably at least 50% sequence identity, more preferably at least 70% sequence identity, still more preferably at least 90% sequence identity. Examples of such kinases are ATR (also known as FRP1, see Cimprich et al, 1996) and DNA-PKcs.

Such molecules may be identified by various means. For instance, information may be obtained about residues which are important for p53 phosphorylation by ATM using alanine scanning and deletion analysis of p53 and/or peptide fragments, for instance the N-terminal 42 amino acids or so of p53, or a fragment of around 10 amino

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acids including the relevant site of phosphorylation. Mutation may be used to identify residues which affect phosphorylation and those which do not. When key residues are identified, computer sequence databases may be scanned for proteins including the same or similar pattern of residues, taking into account conservative variation in sequence (see below) as appropriate. Candidate molecules may then be used in one or more assays for phosphorylation by ATM (such as discussed below).

Identification of key residues for phosphorylation at any of the sites in p53 phosphorylated by ATM may also be used in the design of peptide and non-peptidyl agents which modulate, particularly inhibit, phosphorylation of p53 by ATM, as discussed further below.

Methods of obtaining agents able to modulate interaction between ATM and p53 (or, it must be remembered, ATR, or a related protein having a similar associated kinase activity, and other molecules including a phosphorylation site homologous to one of those phosphorylated in p53 by ATM) include methods wherein a suitable end-point is used to assess interaction in the presence and absence of a test substance. Assay systems may be used to determine ATM kinase activity, ATM DNA binding and/or ATM interaction with one or more other molecules. phosphorylation assays, full-length p53, truncated portions of p53, or portions of p53 fused to other proteins (eg. GST), or a suitable variant or derivative of any of these may be used. Peptide phosphorylation assays may be developed using peptides that correspond to the phosphorylated regions of p53. The phosphorylation of any of the above may be assayed by any of a variety of procedures such as discussed below and may be adapted to high throughput screening approaches. Interference of DNA binding may be assayed but the inhibition of kinase

activity may be more sensitive and identify a greater breadth of inhibitors to DNA binding inhibition, and so may be preferred by the skilled operator of the present invention.

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ATM kinase activity may be assayed for either of the two N-terminal p53 sites. When assaying for phosphorylation, DNA is preferably included in the assay system. Related but different screens may be set up for inhibitors and activators of the two sites of ATM-mediated phosphorylation event.

Generally of most interest is modulation of the phosphorylation of p53 (or other molecule) by ATM.

Detailed disclosure in this respect is included below. It is worth noting, however, that combinatorial library technology provides an efficient way of testing a potentially vast number of different substances for ability to modulate an interaction with and/or activity of a polypeptide. Such libraries and their use are known in the art, for all manner of natural products, small molecules and peptides, among others. The use of peptide libraries may be preferred in certain circumstances.

At the cellular level, A-T cells display chromosomal instability, radiosensitivity, are impaired in p53 induction following treatment with ionising radiation, and show altered regulation of transcription factor NFkB. Thus, the wild-type ATM gene functions as a tumour suppressor, and is a suppressor of neurological degeneration and other degenerative states commonly associated with ageing.

Given the results reported herein on which the present invention is based, activators and inhibitors of ATM-associated kinase activity may be identified and appropriate agents may be obtained, designed and used for

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any of a variety of purposes:

A-T Therapy. Activators of ATM or ATR function may prove to have utility in treating humans with A-T (discussed further below).

Modulation of immune system function. A-T patients display immunodeficiencies, demonstrating that ATM is required for generation of a fully functional immune system. Modulators of ATM or ATR may, therefore, be used in regulating immune system function.

AIDS therapy. It has been shown that the lymphocytes of humans entering the final stages of AIDS have shortened telomeres and this may contribute to them being no longer able to replenish the immune system. Cells of A-T patients lose their telomeres more quickly than those of normal individuals, revealing that ATM plays a positive role in telomere length homeostasis. Activators of ATM function may, therefore, find utility in treatment of individuals with AIDS through lengthening the telomeres of senescent lymphocytes in these individuals, thus allowing replenishment of the immune system.

25 The identification of the site of p53 p53 therapy. phosphorylated by ATM indicates that this of extreme regulatory importance. Indeed, the N-terminal sites on p53 phosphorylated by ATM reside within the region known as "conserved region I" that has been shown to function together with flanking sequences in the interaction with 30 the protein Mdm-2 (see Kussie et al 1996; Picksley et al., 1994; Momand et al., 1992; Chen et al., 1993 and references therein). Mdm-2 serves as a negative regulator of p53 by two mechanisms. First, it masks the p53 transcriptional activation domain, stopping p53 35 activating genes (Momand et al., 1992). Second, Mdm-2 has been shown recently to target p53 for degradation

within the cell (Kubbutat et al., 1997; Haupt et al., 1997). Our data therefore provide an indication that phosphorylation of p53 by ATM will disrupt its interactions with Mdm-2, thus resulting in increased levels of transcriptionally active p53. This knowledge may, therefore, be utilised to generate novel therapeutic agents that target p53 - such as small molecules that, through binding to mutant p53, mimic ATM-mediated activation of this molecule.

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Phosphorylation at any one or more of these sites may affect interaction of p53 with a number of proteins. Mdm2 is one particularly example given the location of Thr18 within the site on p53 to which Mdm2 binds (see e.g. Chen et al., (1993), Kussie et al., (1996), Picksley et al., 15 (1994) and Momand et al., (1992) for characterisation of this interaction) and Ser15 which lies immediately adjacent to the minimal Mdm2 binding sequence. report by Shieh et al published in October 1997 indicates that phosphorylation at Ser15 can disrupt the p53-Mdm2 20 interaction. Phosphorylation of p53 may be used to affect interaction of p53 with any of a number of other proteins, including CBP (Gu et al.; Lill et al.), adenovirus E1B protein, which binds within the amino terminal 123 amino acids of p53 (Kao et al., 1990), with 25 residues Leu-22 and Trp-23 playing an important role (Lin et al., 1994), transcription factors XPD (Rad3) and XPB, as well as CSB involved in strand-specific DNA repair (Wang et al., 1995), TFIIH (Xiao et al., 1994), E2F1 and DP1 (O'Connor et al., 1995), Cellular 30 Replication Protein A (Li and Botchan, 1993), replication factor RPA (Dutta et al., 1993), WT1 (Maheswaran et al., 1993), TATA-binding protein (Seto et al., 1992, Truant et al., 1992, Martin et al., 1993), and TAF(II)40 and 35 TAF(II)60 (Thut et al., 1995).

An assay according to the present invention as discussed

further below may determine the role of phosphorylation of p53 by ATM on any of these interactions and an agent found to be able to modulate such phosphorylation may be used to disrupt or promote any of these interactions, e.g. in a therapeutic context.

Modulating telomere length. A-T cells show accelerated rates of telomere shortening (Metcalfe et al., 1996, Nature Genetics 13, 350-353). Thus, regulators of ATM activity may be used to control telomere length. ATM does not appear to be part of the telomerase enzyme itself (Metcalfe et al. shows that telomerase levels are normal in A-T cells; also, our data and the data of Pandita et al. 1995 show that A-T cells have somewhat shortened telomeres but do not have repressed levels of telomerase). Thus, ATM works not as part of telomerase but as part of a telomere length homeostatic mechanism. It is therefore likely that anti-ATM drugs will work synergistically with anti-telomerase drugs.

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Ageing. A-T patients display enhanced rates of ageing, display a number of symptoms associated with increased age (neurological deterioration, cancers, immunological deficiencies etc), and their cells show shortened lifespan in culture. Agents that modulate ATM activity may therefore be used to treat/prevent disease states associated with premature and normal ageing.

Tumour/Cancer therapy. This is discussed below. Drugs that modulate ATM action may be used to treat A-T patients; treat cancer - through affecting cellular growth capacity by shortening cells telomeres; manipulate the immune system - A-T patients are somewhat immunodeficient; treat cancer - radiosensitization of tumours etc (see below). Also, ATM modulators may be used to limit cell growth potential by affecting telomere length etc. The linkage to p53 may allow p53 therapy,

activating p53 in cancer cells, which may lead to cell growth arrest and/or cell death via apoptosis or another route.

Activators of ATM (or ATR, DNA-PK or related kinases) may 5 be used, for example, to inhibit cell proliferation by activating cell cycle checkpoint arrest in the absence of cellular damage, which may be used in the treatment of tumours, cancer, psoriasis, arteriosclerosis and other hyper-proliferative disorders. Activators may be 10 employed to activate p53 in cells without damaging the cells. Cells of a patient may be treated so that normal cells (p53+) stop growing and are thus refractory to killing by administration of a drug that kills cells via interfering with cell division or DNA replication, while 15 tumour cells (many of which are p53 negative) do not arrest and are consequently selectively killed by the aforementioned agents. By way of example, ATM activators include peptides capable of recognising and binding to both ATM and p53 but which do not interfere with the 20 phosphorylation of the Ser15 and Thr18 sites of p53, or substances capable of activating ATM in a similar manner to the activation observed using DNA.

Cancer radiotherapy and chemotherapy may be augmented using agents in accordance with the present invention. Ionising radiation (IR) and radiomimetic drugs are used commonly to treat cancers, and kill cancer cells predominantly via inflicting DNA damage. Cells deficient in ATM are hypersensitive to ionising radiation and radiomimetics. Thus, inhibitors of the ATM will hypersensitise cells to the killing effects of ionising radiation and radiomimetics. ATM inhibitors may thus be used as adjuncts in cancer radiotherapy and chemotherapy.

Cell growth capacity may be modulated e.g. in treatment of cancer, ageing, and AIDS. It is established that ATM

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plays a crucial role in controlling the length of : télomeric chromosomal ends (Metcalfe et al.). ends in most normal cell types shorten at each cell division, and cells with excessively shortened telomeres are unable to divide. Thus, telomeres are thought to function as a "division counting apparatus" that limits the proliferative capacity of most normal mammalian Inhibitors of ATM function may, therefore, have utility in preventing cancer progression by limiting the growth potential of cancerous or pre-cancerous cells. Activators of ATM may be used to release senescent cells from growth arrest and may thus have utility in treatments of aged individuals. In addition, it has been shown recently that the lymphocytes of humans entering the final stages of AIDS have shortened telomeres and this may contribute to these cells being no longer able to proliferate and replenish the immune system. activators may, therefore, result in lengthening of the telomeres of such cells and restoring their proliferative capacity.

Interaction between ATM and p53 may be inhibited by inhibition of the production of the relevant protein. For instance, production of one or more of these components may be inhibited by using appropriate nucleic acid to influence expression by antisense regulation. The use of anti-sense genes or partial gene sequences to down-regulate gene expression is now well-established. Double-stranded DNA is placed under the control of a promoter in a "reverse orientation" such that transcription of the "anti-sense" strand of the DNA yields RNA which is complementary to normal mRNA transcribed from the "sense" strand of the target gene. The complementary anti-sense RNA sequence is thought then to bind with mRNA to form a duplex, inhibiting translation of the endogenous mRNA from the target gene into protein. Whether or not this is the actual mode of

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action is still uncertain. However, it is established fact that the technique works.

Another possibility is that nucleic acid is used which on transcription produces a ribozyme, able to cut nucleic acid at a specific site - thus also useful in influencing gene expression. Background references for ribozymes include Kashani-Sabet and Scanlon, 1995, Cancer Gene Therapy, 2(3): 213-223, and Mercola and Cohen, 1995, Cancer Gene Therapy, 2(1), 47-59.

Thus, various methods and uses of modulators, which inhibit or potentiate interaction of ATM and p53, particularly phosphorylation of p53 by ATM, are provided as further aspects of the present invention. The purpose of disruption, interference with or modulation of interaction between ATM and p53, particularly the phosphorylation of p53 by ATM may be to modulate any activity mediated by virtue of such interaction, as discussed above and further below.

Various aspects of the present invention relate to modulation of interaction between ATM and DNA. interaction is established here we believe for the first time, and is further shown to have an effect on p53 phosphorylation by ATM. It was surprising that ATM is a DNA binding protein, as there are data suggesting that it is associated with microsomal membranes in the cytoplasm (Watters et al, 1997 and Brown et al, 1997; show ATM is also present in cytoplasmic vesicles) and A-T cells have also been reported to be defective in signalling from the cell membrane in B- and T-cells (see above). It was furthermore surprising that ATM would bind DNA so well. The purification method used and described below does not purify a variety of other (known) DNA binding factors, yet ATM is purified very selectively (about 100-fold in a single step) using a DNA affinity chromatography

procedure.

The present invention provides in one aspect the use of DNA for purifying ATM or ATR. In further aspects, the present invention provides for the use of DNA in assays for activity of ATM or ATR, particularly phosphorylation of p53 (or other molecule).

We have also purified ATM and ATR via another surprising route, using nitrilo-tri-acetic acid (NTA) agarose. 10 has 4 chelating sites for Ni²⁺. Another Ni²⁺ matrix, iminodiacetic acid (IDA) agarose (with 3-chelating sites for Ni²⁺) we have found to bind ATM only weakly. Ni²⁺ matrices are generally used interchangeably to purify proteins that chelate metal ions, usually, via a run of 15 His residues (usually 6 give best binding). ATM does not have a run of 6, 5 or even 4 His residues, so it is surprising that ATM or ATR is purifiable by the Ni-linked columns. Furthermore, since the two matrices are 20 generally used interchangeably, it is further surprising that ATM binds to the NTA well but only poorly to the IDA matrix.

ATM no doubt works in concert with other factors in the detection and signalling of DNA damage. Indeed, although 25 our data reveal that ATM possesses intrinsic DNA-stimulated p53 kinase function, we have observed repeatedly that the presence of additional polypeptides correlates with increased ATM activity. Thus, our most highly purified preparations have considerably less 30 activity than preparations containing an equivalent amount of ATM but also possessing additional co-purifying polypeptides. It is likely that these serve to help tether ATM to the DNA and/or trigger its kinase activity 35 by altering the conformation of the ATM polypeptide. Accordingly, references to ATM, or a protein having a associated kinase activity, include both purified ATM (or

the related protein) and ATM (or the related protein) in combination with associated polypeptides or co-factors present in preparations as obtainable by the methods described herein.

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Assays according to the present invention may be used in the identification of such additional polypeptides, for example by assaying for protein fractions that stimulate ATM activity. The use of ATM or ATR in identifying and/or obtaining cofactors which (e.g. naturally) enhance its kinase activity is further provided by the present invention. ATM activity may under certain circumstances be masked by one or more factors (see discussion section below). Accordingly, the present invention also provides for the use of ATM in identifying and/or obtaining such factors.

Protein or other co-factors of ATM, e.g. which enhance ATM kinase activity, may be used in the design of inhibitors of this, providing another route for modulating ATM activity. This may similarly be used to provide a route to deriving agents that activate ATM, e.g. by inhibiting one or more repressors of ATM activity.

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Brief Description of the Figures
Figure 1: ATM binds to DNA. (A) ATM binds to a dsDNA
oligonucleotide. HeLa nuclear extract was bound to
either streptavidin iron oxide beads (-DNA) or
streptavidin iron oxide beads bearing a 50-mer ds DNA
oligonucleotide (+DNA). After extensive washing, ATM was
eluted from DNA in 500 mM KCl. Eluted proteins were
subjected to 7% SDS-PAGE and ATM visualised by Western
blotting using ATM.B antiserum. (B) Binding of ATM is
dependent on DNA length. ATM enriched extract was bound
to streptavidin iron oxide beads attached to ds DNA of
various sizes (15, 25, 50 or 75 bp). After extensive

washing, ATM was eluted by sequential washes with 100, 250 and 500 mM KCl. Eluates were analysed as in (A). (C) ATM binds DNA containing a variety of different architectures. ATM enriched extract was bound to streptavidin iron oxide beads bound to either ss or ds DNA containing a nick, ds/ss transition, gap or 10 bp insertion. Washing, elution and ATM detection was as in (B).

- Figure 2: Purification of ATM from HeLa cell nuclear 10 extract. (A) ATM Purification strategy. HeLa nuclear extract was subjected to ion exchange chromatography using Q-Sepharose and peak ATM fractions, eluting between 160-200 mM KCl, were passed over heparin-agarose ion 15 exchange resin. ATM fractions eluting from heparinagarose between 200-220 mM KCl were pooled and subjected to DNA affinity purification and elution from DNA-bearing beads at 500 mM KCl resulting in an essentially homogeneous preparation of ATM. (B) Purification of ATM 20 to essential homogeneity. Equivalent volumes (5 μ l) of HeLa cell nuclear extract (50 μ g protein), or pooled fractions following Q-sepharose, Heparin-agarose or DNA affinity chromatography were subjected to 7% SDS-PAGE and proteins visualised by silver staining (upper panel). 25 Fractions were also subjected to Western blot analysis (lower panel) using antibodies raised against ATM, DNA- PK_{cs} , Ku70 plus Ku80 or the 70kDa subunit of RPA, as indicated.
- Figure 3: Purified ATM possesses an associated p53 kinase activity. (A) Analysis of putative ATM substrates. DNA-PK_{cs} (60 ng), Ku (100 ng), Spl (100 ng), p53 (100 ng), RPA-p34 (100 ng) or PCNA (100 ng) were used in kinase reactions in conjunction with approximately 11 fmole of purified ATM (see Experimental Procedures). Proteins were resolved on either 7% (left panel) or 10% (right panel) polyacrylamide gels and phosphorylated

proteins detected by autoradiography. (B) Analysis of total proteins immuno-precipitated from purified ATM preparations. Purified ATM was biotinylated and subjected to immunoprecipitation using either pre-immune sera, or ATM antisera raised against amino acid residues 5 1980-2337 (ATM.B) or the N-terminus (ATM.N) of ATM. Precipitated proteins were resolved on 7.8% polyacrylamide gels and, after transfer to nitrocellulose, total precipitated proteins were detected by probing filters with streptavidin-conjugated 10 horseradish peroxidase. (C) Immunoprecipitated ATM. possesses p53 kinase activity. Purified ATM was immunoprecipitated using pre-immune sera, or anti-ATM antisera ATM.B or ATM.N. Following immunoprecipitation, kinase reactions were performed either in the presence or 15 absence of p53 as indicated. Phosphorylated proteins were resolved on 10% polyacrylamide gels and detected by autoradiography.

20 Figure 4: A DNA-stimulated protein kinase activity copurifies with ATM. (A) ATM associated kinase activity is stimulated by linear DNA containing multiple p53 binding sites. Purified ATM, DNA-PK or cyclin A/cdk2 (11 fmole), as indicated, were used in kinase reactions containing 25 p53 either in the absence (-) or presence of 0.03, 0.3 or 3 fmole of linear DNA bearing multiple p53 binding sites (pG₁₃CAT). Proteins were resolved on 10% polyacrylamide gels and phosphorylated proteins visualised by autoradiography. (B) ATM associated kinase activity does 30 not require DNA ends. In vitro kinase reactions containing 11 fmole of purified ATM in conjunction with p53 were performed in either the absence (-) or presence of 0.03, 0.3 or 30 fmole of linear or supercoiled pG13CAT DNA. Proteins were detected as in (A).

Figure 5: ATM phosphorylates p53 at Ser15 and Thr18 in the presence of DNA. Kinase reactions employing ATM and

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p53 were performed in the presence and absence of DNA. These studies revealed phosphorylation of p53 was increased in the presence of DNA. (A,B) Bands corresponding to 32P-labelled p53 were excised from a gel, digested with trypsin, and chromatographed on a Vydac 5 218TP54 C18 column (see Experimental Procedures). Purified p53 fractions phosphorylated by ATM preparations in the presence, but not in the absence, of DNA (peptides 2a, 2b and 2c) were subjected to peptide sequence analysis as described in Experimental Procedures; 10 radioactivity was measured after each cycle of Edman degradation. The putative amino acid sequence of the p53 peptide showing incorporation of 32P is indicated in panel (D) Tryptic peptide map of p53 phosphorylated by DNA-PK in the presence of DNA. Kinase reactions containing 15 DNA-PK and p53 were performed in the presence of linear DNA and ^{32}P -labelled p53 was analysed as in (A,B), again revealing phosphorylation at Ser15 and Thr18.

- Figure 6a shows the amino acid sequence of human ATM, with the kinase domain marked by underlining. Figure 6b shows the ATM nucleic acid sequence with the initiation codon underlined.
- Figure 7a shows the amino acid sequence of human p53 with residues phosphorylated by ATM marked by underlining. Figure 7b shows the p53 nucleic acid sequence with the initiation codon underlined.
- Figure 8a shows the amino acid sequence of human ATR (FRP-1). Figure 8b shows the ATR nucleic acid sequence with the initiation codon underlined.
- Figure 9a shows the amino acid sequence of DNA-PKcs.

 Figure 9b shows the DNA-PK nucleic acid sequence with the initiation codon underlined.

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Figure 10 shows fractionation of two DNA activated kinase activities in HeLa nuclear cell extract capable of phosphorylating Ser15 of p53. Top panel; a Western immuno-blot was performed with antibodies that specifically recognise p53 phosphorylated on Ser15 on reactions in which fractions generated when HeLa cell nuclear extract was fractionated on Q-sepharose were incubated with p53 and ATP in the presence of sonicated calf thymus DNA. Middle panel; the same set of fractions were tested for DNA-PKcs by using an anti-DNA-PKcs antiserum in western immuno-blot analysis. Lower panel: the same set of fractions were tested for the presence of ATR by using an anti-ATR antiserum in western immuno-blot Additional studies revealed that both activities detected are stimulated by DNA.

Figure 11 shows DNA activated kinase activity (activity 1) co-fractionates with ATR. Activity peak 1 was fractionated further on DNA-cellulose followed by chromatography on Heparin-agarose. Bottom panel; the final set of fractions was tested for p53 kinase activity via incubation with p53, ATP and DNA and then analysis by SDS-polyacrylamide gel electrophoresis and Western immuno-blotting using the p53 Ser15-specific antibodies.

Top panel; a silver-stain of an SDS-polyacrylamide gel of the same set of fractions tested for p53 kinase activity. ATR is indicated with an arrow.

The present invention in various aspects provides for modulating, interfering with or interrupting, increasing or potentiating interaction between the ATM protein and p53, particularly phosphorylation of p53 by ATM, using an appropriate agent. As noted, it having now been established for the first time that ATM is a protein kinase, it is highly likely to act on other molecules, particularly proteins including a site which is homologous to one of the sites in p53 phosphorylated by

ATM. The present invention further extends to the use of proteins having an associated kinase activity similar to ATM, especially DNA-PK and ATR. The present invention extends to modulation of such phosphorylation and this should be borne in mind when considering the disclosure herein which for convenience uses p53 for illustrative purposes, and as a preferred embodiment in certain contexts.

- An agent capable of modulating interaction between ATM and p53 may be capable of blocking interaction between a site located within amino acid residues including Ser15 or Thr18.
 - In addition to interacting at the site of phosphorylation of p53, ATM and p53 may interact at one or more other sites within either or both proteins. Affecting interaction at such a site may have an effect on phosphorylation of p53 by ATM. Various fragments and derivatives of the proteins, particular of p53, may be used to analyse this, using techniques such as alanine scanning and deletion analysis. The present invention encompasses modulation of interaction between ATM and p53 at any site, preferably resulting in modulation of p53 phosphorylation by ATM.

The full amino acid sequence of the ATM protein has been elucidated and is set out in Savitsky et al 1995a, 1995b, and Figure 6a, of which the amino acid residue numbering is used. The kinase domain is marked in Figure 6a. The p53 amino acid sequence is shown in Figure 7a, of which the amino acid residue numbering is used. These sequences are human sequences. ATM and p53 are conserved among vertebrates, particular mammals - see e.g. Figure 2 of Soussi et al. For p53 conservation in the regions of the residues shown herein to be phosphorylated by ATM - so the present invention extends to use in any of its

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aspects of other vertebrate, particularly mammalian, p53 and/or ATM, e.g. primate, such as monkey, rodent, such as mouse or rat, pig, horse, cow, sheep, goat, dog, cat, and so on. The amino acid and nucleic acid sequences of ATR (also known as FRP1) are set out in Cimpich et al, 1996. The amino acid sequence is reproduced as Figure 8a. The amino acid sequence of DNA-PK is provided in Hartley et al, 1995 and is set out in Figure 9a. The nucleic acid sequences of these proteins are also included as Figures 6b, 7b, 8b and 9b.

Agents useful in accordance with the present invention may be identified by screening techniques which involve determining whether an agent under test inhibits or disrupts the interaction of ATM protein or a suitable fragment thereof (e.g. including amino acid residues of the kinase domain, as marked on Figure 6, or a smaller fragment of any of these regions) of ATM, with p53 or a fragment thereof, or a suitable analogue, fragment or variant thereof. One class of preferred fragments of p53 are those which include one or both of the phosphorylation sites at Ser15 or Thr18.

Suitable fragments of ATM or p53 include those which include residues which interact with the counterpart protein. Smaller fragments, and analogues and variants of this fragment may similarly be employed, e.g. as identified using techniques such as deletion analysis or alanine scanning.

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Thus, the present invention provides a peptide fragment of ATM which is able to interact with p53 and/or inhibit interaction between ATM and p53, particularly phosphorylation of p53 by ATM, and provides a peptide fragment of p53 which is able to interact with ATM and/or inhibit interaction between p53 and ATM, particularly phosphorylation of p53 by ATM, such peptide fragments

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being obtainable by means of deletion analysis and/or alanine scanning of the relevant protein - making an appropriate mutation in sequence, bringing together a mutated fragment of one of the proteins with the other or a fragment thereof and determining interaction, preferably phosphorylation of p53 or fragment thereof. In preferred embodiments, the peptide is short, as discussed below, and may be a minimal portion that is able to interact with the relevant counterpart protein and/or inhibit the relevant interaction.

Screening methods and assays are discussed in more detail below.

One class of agents that can be used to disrupt the 15 interaction of ATM and p53 are peptides based on the sequence motifs of ATM or p53 that interact with counterpart p53 or ATM (as discussed already above). Such peptides tend to be short, and may be about 40 amino acids in length or less, preferably about 35 amino acids 20 in length or less, more preferably about 30 amino acids in length, or less, more preferably about 25 amino acids or less, more preferably about 20 amino acids or less, more preferably about 15 amino acids or less, more preferably about 10 amino acids or less, or 9, 8, 7, 6, 5 25 or less in length. The present invention also encompasses peptides which are sequence variants or derivatives of a wild type ATM or p53 sequence, but which retain ability to interact with p53 or ATM (respectively, as the case may be) and/or ability to modulate 30 interaction between ATM and p53, particularly phosphorylation of p53 by ATM.

Instead of using a wild-type ATM or p53 fragment, a

peptide or polypeptide may include an amino acid sequence which differs by one or more amino acid residues from the wild-type amino acid sequence, by one or more of

addition, insertion, deletion and substitution of one or more amino acids. Thus, variants, derivatives, alleles, mutants and homologues, e.g. from other organisms, are included.

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Preferably, the amino acid sequence shares homology with a fragment of the relevant ATM or p53 fragment sequence shown preferably at least about 30%, or 40%, or 50%, or 60%, or 70%, or 75%, or 80%, or 85%, 90% or 95% homology. Thus, a peptide fragment of ATM or p53 may include 1, 2, 3, 4, 5, greater than 5, or greater than 10 amino acid alterations such as substitutions with respect to the wild-type sequence. Preferably, the peptide fragments of ATM are based on the sequence of all or part of the kinase domain as shown in figure 6. Preferably, the p53 fragments are based on the N-terminal sequence of the molecule around the sites phosphorylated by ATM, i.e comprising the amino acid motif PPLSQETFSD, or more generally, the motif SxxT, where x is any amino acid.

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A derivative of a peptide for which the specific sequence is disclosed herein may be in certain embodiments the same length or shorter than the specific peptide. In other embodiments the peptide sequence or a variant thereof may be included in a larger peptide, as discussed above, which may or may not include an additional portion of ATM or p53. 1, 2, 3, 4 or 5 or more additional amino acids, adjacent to the relevant specific peptide fragment in ATM or p53, or heterologous thereto may be included at one end or both ends of the peptide.

(It should not be forgotten that references to ATM and p53 apply equally to ATM and related proteins such as ATR and DNA-PK and other proteins including a phosphorylation site homologous to one in p53 phosphorylated by ATM.)

As is well-understood, homology at the amino acid level

is generally in terms of amino acid similarity or Similarity allows for "conservative identity. variation", i.e. substitution of one hydrophobic residue such as isoleucine, valine, leucine or methionine for another, or the substitution of one polar residue for another, such as arginine for lysine, glutamic for aspartic acid, or glutamine for asparagine. Similarity may be as defined and determined by the TBLASTN program, of Altschul et al. (1990) J. Mol. Biol. 215: 403-10, which is in standard use in the art. Homology may be over the full-length of the relevant peptide or over a contiguous sequence of about 5, 10, 15, 20, 25, 30, 35, 50, 75, 100 or more amino acids, compared with the relevant wild-type amino acid sequence.

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As noted, variant peptide sequences and peptide and nonpeptide analogues and mimetics may be employed, as discussed further below.

Various aspects of the present invention provide a substance, which may be a single molecule or a composition including two or more components, which includes a peptide fragment of ATM or p53 which includes a sequence as recited in Figure 6 or Figure 7, 25 particularly within the ATM kinase domain marked in Figure 6, a peptide consisting essentially of such a sequence, a peptide including a variant, derivative or analogue sequence, or a non-peptide analogue or mimetic which has the ability to interact with ATM or p53 and/or modulate, disrupt or interfere with interaction between 30 ATM or p53.

Variants include peptides in which individual amino acids can be substituted by other amino acids which are closely related as is understood in the art and indicated above.

Non-peptide mimetics of peptides are discussed further

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below.

As noted, a peptide according to the present invention and for use in various aspects of the present invention may include or consist essentially of a fragment of ATM or p53 as disclosed, such as a fragment whose sequence is shown in Figure 6 or Figure 7, respectively. Where one or more additional amino acids are included, such amino acids may be from ATM or p53 or may be heterologous or foreign to ATM or p53. A peptide may also be included within a larger fusion protein, particularly where the peptide is fused to a non-ATM or p53 (i.e. heterologous or foreign) sequence, such as a polypeptide or protein domain.

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The invention also includes derivatives of the peptides, including the peptide linked to a coupling partner, e.g. an effector molecule, a label, a drug, a toxin and/or a carrier or transport molecule, and/or a targeting molecule such as an antibody or binding fragment thereof 20 or other ligand. Techniques for coupling the peptides of the invention to both peptidyl and non-peptidyl coupling partners are well known in the art. In one embodiment, the carrier molecule is a 16 aa peptide sequence derived 25 from the homeodomain of Antennapedia (e.g. as sold under the name "Penetratin"), which can be coupled to a peptide via a terminal Cys residue. The "Penetratin" molecule and its properties are described in WO 91/18981.

Peptides may be generated wholly or partly by chemical synthesis. The compounds of the present invention can be readily prepared according to well-established, standard liquid or, preferably, solid-phase peptide synthesis methods, general descriptions of which are broadly available (see, for example, in J.M. Stewart and J.D. Young, Solid Phase Peptide Synthesis, 2nd edition, Pierce Chemical Company, Rockford, Illinois (1984), in M.

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Bodanzsky and A. Bodanzsky, The Practice of Peptide Synthesis, Springer Verlag, New York (1984); and Applied Biosystems 430A Users Manual, ABI Inc., Foster City, California), or they may be prepared in solution, by the liquid phase method or by any combination of solid-phase, liquid phase and solution chemistry, e.g. by first completing the respective peptide portion and then, if desired and appropriate, after removal of any protecting groups being present, by introduction of the residue X by reaction of the respective carbonic or sulfonic acid or a reactive derivative thereof.

Another convenient way of producing a peptidyl molecule according to the present invention (peptide or polypeptide) is to express nucleic acid encoding it, by use of nucleic acid in an expression system.

Accordingly the present invention also provides in various aspects nucleic acid encoding the polypeptides and peptides of the invention.

Generally, nucleic acid according to the present invention is provided as an isolate, in isolated and/or purified form, or free or substantially free of material with which it is naturally associated, such as free or substantially free of nucleic acid flanking the gene in the human genome, except possibly one or more regulatory sequence(s) for expression. Nucleic acid may be wholly or partially synthetic and may include genomic DNA, cDNA or RNA. Where nucleic acid according to the invention includes RNA, reference to the sequence shown should be construed as reference to the RNA equivalent, with U substituted for T.

Nucleic acid sequences encoding a polypeptide or peptide in accordance with the present invention can be readily prepared by the skilled person using the information and

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references contained herein and techniques known in the art (for example, see Sambrook, Fritsch and Maniatis, "Molecular Cloning, A Laboratory Manual, Cold Spring Harbor Laboratory Press, 1989, and Ausubel et al, Short Protocols in Molecular Biology, John Wiley and Sons, 1992), given the nucleic acid sequence and clones available. These techniques include (i) the use of the polymerase chain reaction (PCR) to amplify samples of such nucleic acid, e.g. from genomic sources, (ii) chemical synthesis, or (iii) preparing cDNA sequences. DNA encoding ATM or p53 fragments may be generated and used in any suitable way known to those of skill in the art, including by taking encoding DNA, identifying suitable restriction enzyme recognition sites either side of the portion to be expressed, and cutting out said portion from the DNA. The portion may then be operably linked to a suitable promoter in a standard commercially available expression system. Another recombinant approach is to amplify the relevant portion of the DNA with suitable PCR primers. Modifications to the ATM or p53 sequences can be made, e.g. using site directed mutagenesis, to lead to the expression of modified ATM or p53 peptide or to take account of codon preference in the host cells used to express the nucleic acid.

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In order to obtain expression of the nucleic acid sequences, the sequences can be incorporated in a vector having one or more control sequences operably linked to the nucleic acid to control its expression. The vectors may include other sequences such as promoters or enhancers to drive the expression of the inserted nucleic acid, nucleic acid sequences so that the polypeptide or peptide is produced as a fusion and/or nucleic acid encoding secretion signals so that the polypeptide produced in the host cell is secreted from the cell. Polypeptide can then be obtained by transforming the

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functional, culturing the host cells so that the polypeptide is produced and recovering the polypeptide from the host cells or the surrounding medium.

Prokaryotic and eukaryotic cells are used for this purpose in the art, including strains of E. coli, yeast, and eukaryotic cells such as COS or CHO cells.

Thus, the present invention also encompasses a method of making a polypeptide or peptide (as disclosed), the method including expression from nucleic acid encoding the polypeptide or peptide (generally nucleic acid according to the invention). This may conveniently be achieved by growing a host cell in culture, containing such a vector, under appropriate conditions which cause or allow expression of the polypeptide. Polypeptides and peptides may also be expressed in in vitro systems, such as reticulocyte lysate.

Systems for cloning and expression of a polypeptide in a variety of different host cells are well known. Suitable host cells include bacteria, eukaryotic cells such as mammalian and yeast, and baculovirus systems. Mammalian cell lines available in the art for expression of a heterologous polypeptide include Chinese hamster ovary cells, HeLa cells, baby hamster kidney cells, COS cells and many others. A common, preferred bacterial host is E. coli.

Suitable vectors can be chosen or constructed, containing appropriate regulatory sequences, including promoter sequences, terminator fragments, polyadenylation sequences, enhancer sequences, marker genes and other sequences as appropriate. Vectors may be plasmids, viral e.g. 'phage, or phagemid, as appropriate. For further details see, for example, Molecular Cloning: a Laboratory Manual: 2nd edition, Sambrook et al., 1989, Cold Spring Harbor Laboratory Press. Many known techniques and

protocols for manipulation of nucleic acid, for example in preparation of nucleic acid constructs, mutagenesis, sequencing, introduction of DNA into cells and gene expression, and analysis of proteins, are described in detail in Current Protocols in Molecular Biology, Ausubel et al. eds., John Wiley & Sons, 1992.

Thus, a further aspect of the present invention provides a host cell containing heterologous nucleic acid as disclosed herein.

The nucleic acid of the invention may be-integrated into the genome (e.g. chromosome) of the host cell.

Integration may be promoted by inclusion of sequences which promote recombination with the genome, in accordance with standard techniques. The nucleic acid may be on an extra-chromosomal vector within the cell, or otherwise identifiably heterologous or foreign to the cell.

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A still further aspect provides a method which includes introducing the nucleic acid into a host cell. The introduction, which may (particularly for in vitro introduction) be generally referred to without limitation as "transformation", may employ any available technique. For eukaryotic cells, suitable techniques may include calcium phosphate transfection, DEAE-Dextran, electroporation, liposome-mediated transfection and transduction using retrovirus or other virus, e.g. vaccinia or, for insect cells, baculovirus. For bacterial cells, suitable techniques may include calcium chloride transformation, electroporation and transfection using bacteriophage. As an alternative, direct injection of the nucleic acid could be employed.

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Marker genes such as antibiotic resistance or sensitivity genes may be used in identifying clones containing

nucleic acid of interest, as is well known in the art.

The introduction may be followed by causing or allowing expression from the nucleic acid, e.g. by culturing host cells (which may include cells actually transformed although more likely the cells will be descendants of the transformed cells) under conditions for expression of the gene, so that the encoded polypeptide (or peptide) is If the polypeptide is expressed coupled to an appropriate signal leader peptide it may be secreted from the cell into the culture medium. Following production by expression, a polypeptide or peptide may be isolated and/or purified from the host cell and/or culture medium, as the case may be, and subsequently used as desired, e.g. in the formulation of a composition which may include one or more additional components, such as a pharmaceutical composition which includes one or more pharmaceutically acceptable excipients, vehicles or carriers (e.g. see below).

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Introduction of nucleic acid encoding a peptidyl molecule according to the present invention may take place in vivo by way of gene therapy, to disrupt or interfere with interaction between ATM or p53

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Thus, a host cell containing nucleic acid according to the present invention, e.g. as a result of introduction of the nucleic acid into the cell or into an ancestor of the cell and/or genetic alteration of the sequence endogenous to the cell or ancestor (which introduction or alteration may take place in vivo or ex vivo), may be comprised (e.g. in the soma) within an organism which is an animal, particularly a mammal, which may be human or non-human, such as rabbit, guinea pig, rat, mouse or other rodent, cat, dog, pig, sheep, goat, cattle or horse, or which is a bird, such as a chicken. Genetically modified or transgenic animals or birds

comprising such a cell are also provided as further aspects of the present invention.

This may have a therapeutic aim. (Gene therapy is discussed below). Also, the presence of a mutant, 5 allele, derivative or variant sequence within cells of an organism, particularly when in place of a homologous endogenous sequence, may allow the organism to be used as a model in testing and/or studying substances which modulate activity of the encoded polypeptide in vitro or 10 are otherwise indicated to be of therapeutic potential. Knock-out mice, for instance, may be used to test for radiosensitivity. Conveniently, however, at least preliminary assays for such substances may be carried out in vitro, that is within host cells or in cell-free 15 systems. Where an effect of a test compound is established on cells in vitro, those cells or cells of the same or similar type may be grafted into an appropriate host animal for in vivo testing.

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For instance, p53 function or activity may be measured in an animal system such as a tumour model, e.g. involving a xenograft, relying on active p53. The animal may be subject to radio- or chemo-therapy and a test substance administered. An augmentation of the reaction in the animal to the radio- or chemo-therapy may be indicative of blocking of ATM phosphorylation of p53.

Suitable screening methods are conventional in the art.

They include techniques such as radioimmunosassay, scintillation proximetry assay and ELISA methods. Suitably either the ATM protein or fragment or p53 or fragment, or an analogue, derivative, variant or functional mimetic thereof, is immobilised whereupon the other is applied in the presence of the agents under test. In a scintillation proximetry assay, a biotinylated protein fragment may be bound to

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streptavidin coated scintillant - impregnated beads (produced by Amersham). Binding of radiolabelled peptide is then measured by determination of radioactivity induced scintillation as the radioactive peptide binds to the immobilized fragment. Agents which intercept this are thus inhibitors of the interaction. Further ways and means of screening for agents which modulate interaction between ATM and p53 are discussed below.

- In one general aspect, the present invention provides an assay method for a substance with ability to modulate, e.g. disrupt or interfere with interaction between ATM and p53, the method including:
- (a) bringing into contact a substance according to the invention including a peptide fragment of ATM, or a protein having an associated kinase activity, or a derivative, variant or analogue thereof as disclosed, a substance including the relevant fragment of p53 or a variant, derivative or analogue thereof.

A test compound which disrupts, reduces, interferes with or wholly or partially abolishes interaction between said substances (e.g. including a ATM fragment and including a p53 fragment), and which may modulate ATM and/or p53 activity, may thus be identified.

Agents which increase or potentiate interaction between the two substances may be identified using conditions which, in the absence of a positively-testing agent, prevent the substances interacting.

Another general aspect of the present invention provides an assay method for a substance able to interact with the relevant region of ATM or p53 as the case may be, the method including:

(a) bringing into contact a substance which includes a peptide fragment of ATM or a protein having an

associated kinase activity which interacts with p53 as disclosed, or which includes a peptide fragment of p53 which interacts with ATM or a protein having an associated kinase activity, or a variant, derivative or analogue of such peptide fragment, as disclosed, and a test compound; and,

- (b) determining interaction between said substance and the test compound.
- A test compound found to interact with the relevant portion of ATM may be tested for ability to modulate, e.g. disrupt or interfere with, ATM interaction with p53 and/or ability to affect p53 and/or ATM activity or other activity mediated by ATM or p53 as discussed already above.

Similarly, a test compound found to interact with the relevant portion of p53 may be tested for ability to modulate, e.g. disrupt or interfere with, p53 interaction with ATM and/or ability to affect ATM and/or p53 activity or other activity mediated by p53 or ATM as discussed elsewhere herein.

Another general aspect of the present invention provides an assay method for a substance able to affect p53 activity, the method including:

- (a) bringing into contact p53 and a test compound;and,
 - (b) determining p53 activity.

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p53 activity may be determined in the presence and absence of ATM to allow for an effect of a test compound on activity to be attributed to an effect on interaction between p53 and ATM, preferably phosphorylation of p53 by ATM (discussed further below).

p53 activities which may be determined include induction

of expression of a protein such as p21 (WAF1), cellular sensitivity to ionizing radiation, p53-induced apoptosis activity, p53-induced anti-proliferative activity, p53-induced senescence of cells

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In assaying for agents able to modulate phosphorylation of p53 by ATM, suitable fragments of p53 may be employed including any of the sites of such phosphorylation. Where it is desired to determine phosphorylation at the Ser15 and/or Thr18 site, DNA will generally be included in the assay system to stimulate the requisite kinase activity of ATM. As noted, the present invention extends also to non-human p53 and phosphorylation at sites equivalent to those of human p53 identified herein. Thus, the assays may employ derivatives of full length p53 or the p53 fragments including the phosphorylation sites at Ser15 and/or Thr18.

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The present invention further provides the use of DNA for stimulating phosphorylation of p53 by ATM, e.g. in an assay but also in many other contexts. Such phosphorylation may include at the Ser15 and/or Thr18 site of human p53 or equivalent site in p53 of another species, particularly of a vertebrate such as a mammal.

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An assay according to the present invention may include an inhibitor of DNA-PKcs kinase activity, to avoid complications of redundant phosphorylation by that kinase. Such an inhibitor of DNA-PKcs kinase activity might not affect ATM kinase activity.

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Further assays according to the present invention are for agents which modulate DNA binding by ATM. Inhibitors and/or activators may be screened using appropriate conditions for determination of DNA binding by ATM.

Thus, a further aspect of the present invention provides

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an assay method for a compound able to affect DNA binding by ATM or a protein having an associated kinase activity, the method including:

- (a) bringing into contact a substance which is ATM or a protein having an associated kinase activity, or a fragment, variant or derivative thereof able to bind DNA, DNA and a test compound, under conditions wherein, in the absence of the test compound being an inhibitor of DNA binding by ATM or the protein having an associated kinase activity, said substance binds said DNA; and,
- (b) determining binding between said substance and said DNA.

Activators of DNA binding by ATM may similarly be
identified using an assay method wherein said substance,
the DNA and the test compound are brought together under
conditions wherein in the absence of the test compound
being a potentiator of DNA binding by ATM, the substance
does not bind the DNA. Activators include substances
which activate ATM associated kinase activity in the
absence of DNA or substances which enhance the
interaction of ATM and p53, both of which may allow the
induction of a p53 response in the absence of DNA damage,
e.g. as caused by irradiation.

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DNA binding may be determined using any suitable technique, including an electrophoretic mobility shift assay (EMSA), UV protein-DNA crosslinking, chemical or DNaseI footprinting, and so on.

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Determination of DNA binding by ATM may be performed in conjunction with determination of phosphorylation, sequentially or simultaneously. For instance a preliminary screen may identify molecules which modulate DNA binding by ATM and such substances may then be used in assays to determine their ability (or not) to modulate phosphorylation of p53. The converse, in which ability

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to modulate phosphorylation is determined prior to ability to modulate ATM DNA binding, is also possible, as is to run two assays in parallel.

Preliminary assays in vitro may be followed by, or run in parallel with, in vivo assays.

Of course, the person skilled in the art will design any appropriate control experiments with which to compare results obtained in test assays.

Performance of an assay method according_to the present invention may be followed by isolation and/or manufacture and/or use of a compound, substance or molecule which tests positive for ability to modulate interaction between ATM and p53 and/or inhibit ATM or p53 activity or a mediated activity.

The precise format of an assay of the invention may be varied by those of skill in the art using routine skill 20 and knowledge. For example, interaction between substances may be studied in vitro by labelling one with a detectable label and bringing it into contact with the other which has been immobilised on a solid support. Suitable detectable labels, especially for peptidyl 25 substances include 35S-methionine which may be incorporated into recombinantly produced peptides and polypeptides. Recombinantly produced peptides and polypeptides may also be expressed as a fusion protein 30 containing an epitope which can be labelled with an antibody.

The protein which is immobilized on a solid support may be immobilized using an antibody against that protein bound to a solid support or via other technologies which are known per se. A preferred in vitro interaction may utilise a fusion protein including glutathione-S-

transferase (GST). This may be immobilized on glutathione agarose beads. In an in vitro assay format of the type described above a test compound can be assayed by determining its ability to diminish the amount of labelled peptide or polypeptide which binds to the immobilized GST-fusion polypeptide. This may be determined by fractionating the glutathione-agarose beads by SDS-polyacrylamide gel electrophoresis. Alternatively, the beads may be rinsed to remove unbound protein and the amount of protein which has bound can be determined by counting the amount of label present in, for example, a suitable scintillation counter.

An assay according to the present invention may also take
the form of an in vivo assay. The in vivo assay may be
performed in a cell line such as a yeast strain or
mammalian cell line in which the relevant polypeptides or
peptides are expressed from one or more vectors
introduced into the cell.

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The ability of a test compound to modulate interaction between ATM and p53 may be determined using a so-called two-hybrid assay.

For example, a polypeptide or peptide containing a 25 fragment of ATM or p53 as the case may be, or a peptidyl analogue or variant thereof as disclosed, may be fused to a DNA binding domain such as that of the yeast transcription factor GAL4. (A particularly preferred fragment of ATM may include or be the kinase domain or a 30 fragment of the kinase domain.) The GAL4 transcription factor includes two functional domains. These domains are the DNA binding domain (GAL4DBD) and the GAL4 transcriptional activation domain (GAL4TAD). By fusing one polypeptide or peptide to one of those domains and 35 another polypeptide or peptide to the respective counterpart, a functional GAL4 transcription factor is

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restored only when two polypeptides or peptides of interest interact. Thus, interaction of the polypeptides or peptides may be measured by the use of a reporter gene probably linked to a GAL4 DNA binding site which is capable of activating transcription of said reporter gene. This assay format is described by Fields and Song, 1989, Nature 340; 245-246. This type of assay format can be used in both mammalian cells and in yeast. Other combinations of DNA binding domain and transcriptional activation domain are available in the art and may be preferred, such as the LexA DNA binding domain and the VP60 transcriptional activation domain.

When looking for peptides or other substances which interfere with interaction between a ATM polypeptide or 15 peptide and p53 polypeptide or peptide, the ATM or p53 polypeptide or peptide may be employed as a fusion with (e.g.) the LexA DNA binding domain, and the counterpart p53 or ATM polypeptide or peptide as a fusion with (e.g.) VP60, and involves a third expression cassette, which may 20 be on a separate expression vector, from which a peptide or a library of peptides of diverse and/or random sequence may be expressed. A reduction in reporter gene expression (e.g. in the case of β -galactosidase a weakening of the blue colour) results from the presence 25 of a peptide which disrupts the ATM/p53 interaction, which interaction is required for transcriptional activation of the eta-galactosidase gene. Where a test substance is not peptidyl and may not be expressed from encoding nucleic acid within a said third expression 30 cassette, a similar system may be employed with the test substance supplied exogenously.

When performing a two hybrid assay to look for substances
which interfere with the interaction between two
polypeptides or peptides it may be preferred to use
mammalian cells instead of yeast cells. The same

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principles apply and appropriate methods are well known to those skilled in the art.

In preferred assays according to the present invention, the end-point of the assay, that is to say that which is determined in order to assess the effect of the test agent on the interaction of interest, is phosphorylation of p53 or a fragment, variant or derivative thereof, or other molecule including a phosphorylation site homologous to one of those in p53 phosphorylated by ATM.

Thus, a further aspect of the present invention provides an assay method including

- (a) bringing into contact a substance which includes at least a fragment of ATM which phosphorylates p53, a substance which includes at least a fragment of p53 including a site phosphorylated by ATM, and a test compound; and,
- (b) determining phosphorylation at said site.
 Of course, any suitable variant or derivative of ATM and/or p53 may be employed in such an assay.

Phosphorylation may be determined for example by immobilising p53 or a fragment, variant or derivative thereof, e.g. on a bead or plate, and detecting phosphorylation using an antibody or other binding molecule (such as Mdm2 or a fragment thereof) which binds the relevant site of phosphorylation with a different affinity when the site is phosphorylated from when the site is not phosphorylated. Such antibodies may be obtained by means of any standard technique as discussed elsewhere herein, e.g. using a phosphorylated peptide (such as a fragment of p53). Binding of a binding molecule which discriminates between the phosphorylated and non-phosphorylated form of p53 or relevant fragment, variant or derivative thereof may be assessed using any technique available to those skilled in the art, which

may involve determination of the presence of a suitable label, such as fluorescence. Phosphorylation may be determined by immobilisation of p53 or a fragment, variant or derivative thereof, on a suitable substrate such as a bead or plate, wherein the substrate is impregnated with scintillant, such as in a standard scintillation proximetry assay, with phosphorylation being determined via measurement of the incorporation of radioactive phosphate. Phosphate incorporation into p53 or a fragment, variant or derivative thereof, may be determined by precipitation with acid, such as trichloroacetic acid, and collection of the precipitate on a nitrocellulose filter paper, followed by measurement of incorporation of radiolabeled phosphate.

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An agent able to inhibit phosphorylation of p53 by ATM may include an ATP analogue or other substance able to affect the catalytic properties of the enzymically active site of ATM. An inhibitor of phosphorylation may interact with ATM within the kinase domain marked (for human ATM) in Figure 6. Residues within this domain are involved with interaction with p53 and catalysis of the phosphorylation. Residues outside of the domain may also be involved in interacting with p53 and agents which interfere with such interaction may affect the phosphorylation as discussed elsewhere herein.

The amount of test substance or compound which may be added to an assay of the invention will normally be determined by trial and error depending upon the type of compound used. Typically, from about 0.001 nM to 1mM or more concentrations of putative inhibitor compound may be used, for example from 0.01 nM to $100\mu\text{M}$, e.g. 0.1 to 50 μM , such as about 10 μM . Greater concentrations may be used when a peptide is the test substance. Even a molecule which has a weak effect may be a useful lead compound for further investigation and development.

Compounds which may be used may be natural or synthetic chemical compounds used in drug screening programmes. Extracts of plants which contain several characterised or uncharacterised components may also be used.

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Antibodies directed to the site of interaction in either protein form a further class of putative inhibitor compounds. Candidate inhibitor antibodies may be characterised and their binding regions determined to provide single chain antibodies and fragments thereof which are responsible for disrupting the interaction.

Antibodies may be obtained using techniques which are standard in the art. Methods of producing antibodies include immunising a mammal (e.g. mouse, rat, rabbit, 15 horse, goat, sheep or monkey) with the protein or a fragment thereof. Antibodies may be obtained from immunised animals using any of a variety of techniques known in the art, and screened, preferably using binding of antibody to antigen of interest. For instance, 20 Western blotting techniques or immunoprecipitation may be used (Armitage et al., 1992, Nature 357: 80-82). Isolation of antibodies and/or antibody-producing cells from an animal may be accompanied by a step of 25 sacrificing the animal.

As an alternative or supplement to immunising a mammal with a peptide, an antibody specific for a protein may be obtained from a recombinantly produced library of expressed immunoglobulin variable domains, e.g. using lambda bacteriophage or filamentous bacteriophage which display functional immunoglobulin binding domains on their surfaces; for instance see WO92/01047. The library may be naive, that is constructed from sequences obtained from an organism which has not been immunised with any of the proteins (or fragments), or may be one constructed using sequences obtained from an organism which has been

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exposed to the antigen of interest.

Antibodies according to the present invention may be modified in a number of ways. Indeed the term "antibody" should be construed as covering any binding substance having a binding domain with the required specificity. Thus the invention covers antibody fragments, derivatives, functional equivalents and homologues of antibodies, including synthetic molecules and molecules whose shape mimicks that of an antibody enabling it to bind an antigen or epitope.

Example antibody fragments, capable of binding an antigen or other binding partner are the Fab fragment consisting of the VL, VH, Cl and CH1 domains; the Fd fragment consisting of the VH and CH1 domains; the Fv fragment consisting of the VL and VH domains of a single arm of an antibody; the dAb fragment which consists of a VH domain; isolated CDR regions and F(ab')2 fragments, a bivalent fragment including two Fab fragments linked by a disulphide bridge at the hinge region. Single chain Fv fragments are also included.

A hybridoma producing a monoclonal antibody according to the present invention may be subject to genetic mutation 25 or other changes. It will further be understood by those skilled in the art that a monoclonal antibody can be subjected to the techniques of recombinant DNA technology to produce other antibodies or chimeric molecules which retain the specificity of the original antibody. 30 techniques may involve introducing DNA encoding the immunoglobulin variable region, or the complementarity determining regions (CDRs), of an antibody to the constant regions, or constant regions plus framework 35 regions, of a different immunoglobulin. instance, EP184187A, GB 2188638A or EP-A-0239400. Cloning and expression of chimeric antibodies are

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described in EP-A-0120694 and EP-A-0125023.

Hybridomas capable of producing antibody with desired binding characteristics are within the scope of the present invention, as are host cells, eukaryotic or prokaryotic, containing nucleic acid encoding antibodies (including antibody fragments) and capable of their expression. The invention also provides methods of production of the antibodies including growing a cell capable of producing the antibody under conditions in which the antibody is produced, and preferably secreted.

The reactivities of antibodies on a sample may be determined by any appropriate means. Tagging with individual reporter molecules is one possibility. 15 reporter molecules may directly or indirectly generate detectable, and preferably measurable, signals. linkage of reporter molecules may be directly or indirectly, covalently, e.g. via a peptide bond or noncovalently. Linkage via a peptide bond may be as a 20 result of recombinant expression of a gene fusion encoding antibody and reporter molecule. determining binding is not a feature of the present invention and those skilled in the art are able to choose a suitable mode according to their preference and general 25 knowledge.

Antibodies may also be used in purifying and/or isolating a polypeptide or peptide according to the present invention, for instance following production of the polypeptide or peptide by expression from encoding nucleic acid therefor. Antibodies may be useful in a therapeutic context (which may include prophylaxis) to disrupt the ATM/p53 (or ATR/p53) interaction with a view to inhibiting their activity. Antibodies can for instance be micro-injected into cells, e.g. at a tumour site, subject to radio- and/or chemo-therapy (as

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discussed already above). Antibodies may be employed in accordance with the present invention for other therapeutic and non-therapeutic purposes which are discussed elsewhere herein.

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Other candidate inhibitor compounds may be based on modelling the 3-dimensional structure of a polypeptide or peptide fragment and using rational drug design to provide potential inhibitor compounds with particular molecular shape, size and charge characteristics.

A compound found to have the ability to affect ATM and/or . p53 activity has therapeutic and other potential in a number of contexts, as discussed. For therapeutic treatment such a compound may be used in combination with 15 any other active substance, e.g. for anti-tumour therapy another anti-tumour compound or therapy, such as radiotherapy or chemotherapy. In such a case, the assay of the invention, when conducted in vivo, need not 20 measure the degree of modulation of interaction between p53 and ATM (or appropriate fragment, variant or derivative thereof) or of modulation of p53 phosphorylation or activity caused by the compound being Instead the effect on DNA repair, homologous 25 recombination, cell viability, cell killing (e.g. in the presence and absence of radio- and/or chemo-therapy), retroviral integration, and so on, may be measured. may be that such a modified assay is run in parallel with or subsequent to the main assay of the invention in order 30 to confirm that any such effect is as a result of the inhibition of interaction between ATM and p53 caused by said inhibitor compound and not merely a general toxic effect.

Thus, an agent identified using one or more primary screens (e.g. in a cell-free system) as having ability to interact with ATM and/or p53 and/or modulate activity of

ATM and/or p53 may be assessed further using one or more secondary screens. A secondary screen may involve testing for cellular radiosensitisation and/or sensitisation to radiomimetic drugs, effect on chromosome telomere length, inducing or preventing cell-cycle arrest following irradiation or other cellular insult, an effect of p53 induction following ionising radiation or other cellular insult, or induction of p21 or other downstream p53 target.

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Following identification of a substance or agent which modulates or affects ATM and/or p53 activity, the substance or agent may be investigated further. Furthermore, it may be manufactured and/or used in preparation, i.e. manufacture or formulation, of a composition such as a medicament, pharmaceutical composition or drug. These may be administered to individuals, e.g. for any of the purposes discussed elsewhere herein.

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As noted, the agent may be peptidyl, e.g. a peptide which includes a sequence as recited above, or may be a functional analogue of such a peptide.

- As used herein, the expression "functional analogue" relates to peptide variants or organic compounds having the same functional activity as the peptide in question, which may interfere with the interaction between ATM and p53. Examples of such analogues include chemical compounds which are modelled to resemble the three dimensional structure of the ATM or p53 domain in the contact area, and in particular the arrangement of the
- In a further aspect, the present invention provides the use of the above substances in methods of designing or screening for mimetics of the substances.

key amino acid residues as they appear in ATM or p53.

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Accordingly, the present invention provides a method of designing mimetics of ATM or p53 having the biological activity of p53 or ATM binding or inhibition, the activity of allosteric inhibition of p53 or ATM and/or the activity of modulating, e.g. inhibiting, ATM/p53 interaction, said method comprising:

- (i) analysing a substance having the biological activity to determine the amino acid residues essential and important for the activity to define a pharmacophore; and,
- (ii) modelling the pharmacophore to design and/or screen candidate mimetics having the biological activity.

Suitable modelling techniques are known in the art. This includes the design of so-called "mimetics" which involves the study of the functional interactions fluorogenic oligonucleotide the molecules and the design of compounds which contain functional groups arranged in such a manner that they could reproduced those interactions.

The designing of mimetics to a known pharmaceutically active compound is a known approach to the development of pharmaceuticals based on a "lead" compound. This might be desirable where the active compound is difficult or expensive to synthesise or where it is unsuitable for a particular method of administration, e.g. peptides are not well suited as active agents for oral compositions as they tend to be quickly degraded by proteases in the alimentary canal. Mimetic design, synthesis and testing may be used to avoid randomly screening large number of molecules for a target property.

There are several steps commonly taken in the design of a mimetic from a compound having a given target property. Firstly, the particular parts of the compound that are critical and/or important in determining the target

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property are determined. In the case of a peptide, this can be done by systematically varying the amino acid residues in the peptide, e.g. by substituting each residue in turn. These parts or residues constituting the active region of the compound are known as its "pharmacophore".

Once the pharmacophore has been found, its structure is modelled to according its physical properties, e.g.

stereochemistry, bonding, size and/or charge, using data from a range of sources, e.g. spectroscopic techniques, X-ray diffraction data and NMR. Computational analysis, similarity mapping (which models the charge and/or volume of a pharmacophore, rather than the bonding between atoms) and other techniques can be used in this modelling process.

In a variant of this approach, the three-dimensional structure of the ligand and its binding partner are modelled. This can be especially useful where the ligand and/or binding partner change conformation on binding, allowing the model to take account of this the design of the mimetic.

25 A template molecule is then selected onto which chemical groups which mimic the pharmacophore can be grafted. The template molecule and the chemical groups grafted on to it can conveniently be selected so that the mimetic is easy to synthesise, is likely to be pharmacologically acceptable, and does not degrade in vivo, while retaining the biological activity of the lead compound. The mimetic or mimetics found by this approach can then be screened to see whether they have the target property, or to what extent they exhibit it. Further optimisation or modification can then be carried out to arrive at one or more final mimetics for in vivo or clinical testing.

The mimetic or mimetics found by this approach can then be screened to see whether they have the target property, or to what extent they exhibit it. Further optimisation or modification can then be carried out to arrive at one or more final mimetics for *in vivo* or clinical testing.

Mimetics of this type together with their use in therapy form a further aspect of the invention.

The present invention further provides the use of a peptide which includes a sequence as disclosed, or a derivative, active portion, analogue, variant or mimetic, thereof able to interact with ATM or p53 and/or modulate, e.g. inhibit, interaction between ATM and p53 and/or modulate, e.g inhibit, ATM and/or p53 activity, in screening for a substance able to interact with p53 and/or ATM, and/or modulate, e.g. inhibit, interaction between ATM and p53, and/or inhibit ATM and/or p53 activity.

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Generally, such a substance, e.g. inhibitor, according to the present invention is provided in an isolated and/or purified form, i.e. substantially pure. This may include being in a composition where it represents at least about 90% active ingredient, more preferably at least about 95%, more preferably at least about 98%. Such a composition may, however, include inert carrier materials or other pharmaceutically and physiologicaly acceptable excipients. As noted below, a composition according to the present invention may include in addition to an inhibitor compound as disclosed, one or more other molecules of therapeutic use, such as an anti-tumour agent.

The present invention extends in various aspects not only to a substance identified as a modulator of ATM and p53 interaction and/or ATM or p53-mediated activity, property

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or pathway, in accordance with what is disclosed herein, but also a pharmaceutical composition, medicament, drug or other composition comprising such a substance, a method comprising administration of such a composition to a patient, e.g. for a purpose discussed elsewhere herein, which may include preventative treatment, use of such a substance in manufacture of a composition for administration, e.g. for a purpose discussed elsewhere herein, and a method of making a pharmaceutical composition comprising admixing such a substance with a pharmaceutically acceptable excipient, vehicle or carrier, and optionally other ingredients.

A substance according to the present invention such as an inhibitor of ATM and p53 interaction may be provided for use in a method of treatment of the human or animal body by therapy which affects an ATM or p53-mediated activity in cells, e.g. tumour cells. Other purposes of a method of treatment employing a substance in accordance with the present invention are dicussed elsewhere herein.

Thus, the invention further provides a method of modulating an ATM and/or p53-mediated activity, e.g. for a purpose discussed elsewhere herein, which includes administering an agent which modulates, inhibits or blocks the interaction of ATM with p53 protein, such a method being useful in treatment where such modulation, inhibition or blocking is desirable, or an agent which increase, potentiates or strengthens interaction of ATM with p53, useful in treatment where this is desirable.

The invention further provides a method of treatment which includes administering to a patient an agent which interferes with the interaction of ATM with p53.

Exemplary purposes of such treatment are discussed elsewhere herein.

Whether it is a polypeptide, antibody, peptide, nucleic acid molecule, small molecule, mimetic or other pharmaceutically useful compound according to the present invention that is to be given to an individual, administration is preferably in a "prophylactically effective amount" or a "therapeutically effective amount" (as the case may be, although prophylaxis may be considered therapy), this being sufficient to show benefit to the individual. The actual amount administered, and rate and time-course of administration, will depend on the nature and severity of what is being treated. Prescription of treatment, e.g. decisions on dosage etc, is within the responsibility of general practioners and other medical doctors.

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A composition may be administered alone or in combination with other treatments, either simultaneously or sequentially dependent upon the condition to be treated.

20 Pharmaceutical compositions according to the present invention, and for use in accordance with the present invention, may include, in addition to active ingredient, a pharmaceutically acceptable excipient, carrier, buffer, stabiliser or other materials well known to those skilled in the art. Such materials should be non-toxic and should not interfere with the efficacy of the active ingredient. The precise nature of the carrier or other material will depend on the route of administration, which may be oral, or by injection, e.g. cutaneous, subcutaneous or intravenous.

Pharmaceutical compositions for oral administration may be in tablet, capsule, powder or liquid form. A tablet may include a solid carrier such as gelatin or an adjuvant. Liquid pharmaceutical compositions generally include a liquid carrier such as water, petroleum, animal or vegetable oils, mineral oil or synthetic oil.

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Physiological saline solution, dextrose or other saccharide solution or glycols such as ethylene glycol, propylene glycol or polyethylene glycol may be included.

For intravenous, cutaneous or subcutaneous injection, or injection at the site of affliction, the active ingredient will be in the form of a parenterally acceptable aqueous solution which is pyrogen-free and has suitable pH, isotonicity and stability. Those of relevant skill in the art are well able to prepare suitable solutions using, for example, isotonic vehicles such as Sodium Chloride Injection, Ringer's Injection, Lactated Ringer's Injection. Preservatives, stabilisers, buffers, antioxidants and/or other additives may be included, as required.

Liposomes, particularly cationic liposomes, may be used in carrier formulations.

Examples of techniques and protocols mentioned above can be found in Remington's Pharmaceutical Sciences, 16th edition, Osol, A. (ed), 1980.

The agent may be administered in a localised manner to a tumour site or other desired site or may be delivered in a manner in which it targets tumour or other cells.

Targeting therapies may be used to deliver the active agent more specifically to certain types of cell, by the use of targeting systems such as antibody or cell specific ligands. Targeting may be desirable for a variety of reasons, for example if the agent is unacceptably toxic, or if it would otherwise require too high a dosage, or if it would not otherwise be able to enter the target cells.

Instead of administering these agents directly, they may

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be produced in the target cells by expression from an encoding gene introduced into the cells, eg in a viral vector (a variant of the VDEPT technique - see below). The vector may targeted to the specific cells to be treated, or it may contain regulatory elements which are switched on more or less selectively by the target cells.

The agent (e.g. small molecule, mimetic) may be administered in a precursor form, for conversion to the active form by an activating agent produced in, or targeted to, the cells to be treated. This type of approach is sometimes known as ADEPT or WDEPT, the former involving targeting the activator to the cells by conjugation to a cell-specific antibody, while the latter involves producing the activator, e.g. an enzyme, in a vector by expression from encoding DNA in a viral vector (see for example, EP-A-415731 and WO 90/07936).

An agent may be administered in a form which is inactive but which is converted to an active form in the body. For instance, the agent may be phosphorylated (e.g. to improve solubility) with the phosphate being cleaved to provide an active form of the agent in the body.

A composition may be administered alone or in combination with other treatments, either simultaneously or sequentially dependent upon the condition to be treated, such as cancer, virus infection or any other condition in which a ATM or p53-mediated effect is desirable.

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Nucleic acid according to the present invention, encoding a polypeptide or peptide able to modulate, e.g. interfere with, ATM and p53 interaction and/or induce or modulate activity or other ATM or p53-mediated cellular pathway or function, may be used in methods of gene therapy, for instance in treatment of individuals, e.g. with the aim of preventing or curing (wholly or partially) a disorder

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or for another purpose as discussed elsewhere herein.

Vectors such as viral vectors have been used in the prior art to introduce nucleic acid into a wide variety of different target cells. Typically the vectors are exposed to the target cells so that transfection can take place in a sufficient proportion of the cells to provide a useful therapeutic or prophylactic effect from the expression of the desired polypeptide. The transfected nucleic acid may be permanently incorporated into the genome of each of the targeted cells, providing long lasting effect, or alternatively the treatment may have to be repeated periodically.

A variety of vectors, both viral vectors and plasmid vectors, are known in the art, see US Patent No. 5,252,479 and WO 93/07282. In particular, a number of viruses have been used as gene transfer vectors, including papovaviruses, such as SV40, vaccinia virus, herpesviruses, including HSV and EBV, and retroviruses. Many gene therapy protocols in the prior art have used disabled murine retroviruses.

As an alternative to the use of viral vectors other known methods of introducing nucleic acid into cells includes electroporation, calcium phosphate co-precipitation, mechanical techniques such as microinjection, transfer mediated by liposomes and direct DNA uptake and receptor-mediated DNA transfer.

Receptor-mediated gene transfer, in which the nucleic acid is linked to a protein ligand via polylysine, with the ligand being specific for a receptor present on the surface of the target cells, is an example of a technique for specifically targeting nucleic acid to particular cells.

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A polypeptide, peptide or other substance able to modlate or interfere with the interaction of the relevant polypeptide, peptide or other substance as disclosed herein, or a nucleic acid molecule encoding a peptidyl such molecule, may be provided in a kit, e.g. sealed in a suitable container which protects its contents from the external environment. Such a kit may include instructions for use.

- In further aspects the present invention provides for the 10 provision of purified ATM and purified ATR. Purified ATM or ATR, for instance about 10% pure, more preferably about 20% pure, more preferably about 30% pure, more preferably about 40% pure, more preferably about 50% pure, more preferably about 60% pure, more preferably 15 about 70% pure, more preferably about 80% pure, more preferably about 90% pure, more preferably about 95% pure, or substantially pure ATM or ATR is obtainable using DNA. Such DNA may be in any form which ATM or ATR 20 bind, including single-stranded DNA, double-stranded DNA, nicked DNA, covalently closed DNA circles and so on. is surprising that any and all of these are bound by ATM as shown experimentally below.
- In one aspect the present invention provides the use of DNA for purifying ATM or ATR.

In another aspect the present invention provides a method of purifying ATM or ATR, the method including contacting

ATM or ATR with DNA. A mixture of material including ATM or ATR may be contacted against immobilised DNA (e.g. on a bead or agarose, and either covalently or non-covalently such as via a specific binding molecule such as streptavidin or biotin) and molecules which do not bind washed off.

We have also established that ATM and ATR may be purified

using NTA, preferably in the presence of Ni²⁺. The NTA may be on any suitable support such as agarose or sepharose. Thus, a further aspect of the present invention provides the use of NTA, preferably with Ni²⁺, for purifying ATM or ATR.

Another aspect of the present invention provides a method of purifying ATM or ATR which includes, contacting ATM or ATR with NTA, preferably with Ni²⁺ and washing off molecules which do not bind.

Purification using DNA may be combined with purification using NTA, preferably with Ni^{2+} , sequentially or simultaneously.

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Either technique may be used for identification of cofactors of ATM which modulate ATM activity, such as factors which affect the interaction between ATM and DNA.

The ATM contacted by DNA and/or NTA in a purification may be in a mixture of molecules, such as a cellular extract, such as from a cell of an A-T patient, a normal cell of an organism such as a human or a recombinant host cell expressing the protein from encoding DNA, such as a bacterial, eukaryotic (e.g. mammalian or yeast) or insect cell, such as in a baculovirus expression system. Purification may follow production of ATM recombinantly in a suitable expression system, such as a cell, by expression from encoding nucleic acid.

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Following purification, ATM may be used as desired, e.g. in an assay for an agent which modulates its phosphorylation of p53 or other molecule, in raising or obtaining a specific antibody or other binding molecule, or in a therapeutic context such as to compensate in an individual for the absence of wild-type ATM (as in, for example, a patient with A-T).

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Various further aspects and embodiments of the present invention will be apparent to those skilled in the art in view of the present disclosure. Certain aspects and embodiments of the invention will now be illustrated by way of example and with reference to the figures discussed already above.

ATM binds to DNA

A biotinylated random ds 50-mer oligonucleotide was coupled to streptavidin iron-oxide particles and these were employed to recover DNA binding proteins from HeLa cell nuclear extracts. This approach revealed that ATM interacts with particles bearing this random piece of ds DNA (Figure 1A). This binding is due to the presence of DNA, since streptavidin iron-oxide particles alone are unable to bind ATM (Figure 1A). Importantly, the sequence specific DNA-binding protein Spl and the nonspecific DNA interacting protein complex containing RNA polymerase II (Pol II) are both unable to interact stably with the random DNA fragment employed in these studies (Figure 1A). Furthermore, $\text{DNA-PK}_{\text{CS}}$ present in the crude nuclear extract binds only very inefficiently to the immobilised DNA despite the fact that its DNA-targeting component Ku is present (data not shown). Notably, protein quantification reveals that, under conditions in which over 90% of ATM binds to the DNA-coupled particles, less than 2% of total nuclear protein is retained. Hence, the retention of ATM by DNA in these studies is highly specific.

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The above assay revealed that ATM, or an ATM complex, is capable of binding to a random piece of duplex DNA. Additional studies revealed that ATM is also retained by particles containing another unrelated oligonucleotide, suggesting strongly that the interaction is not sequence-specific (data not shown). To investigate the DNA binding properties of ATM further, we tested a series of

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DNAs with a variety of sizes and architectures. In these studies, binding and initial washes were conducted in the presence of 50 mM KC1, then bound material was eluted by sequential washes at 100, 250 and 500 nM KC1. Figure 1B demonstrates that the interaction between DNA and ATM is dependent on the size of the DNA-duplex. Thus, with a ds 15-mer, some ATM is still present in the unbound fraction and most bound material elutes in the lower salt wash. However, as the duplex size is increased, it becomes progressively more effective at binding ATM, such that when ds oligonucleotides of 50 bp or larger are employed, binding of ATM is almost quantitative and all bound ATM elutes in the higher salt wash (Figure 1B).

Since a variety of DNA structures are known to be 15 produced by IR and are present during DNA-repair processes, we assessed the ability of ATM to bind to various types of DNA structure. Thus, assays were conducted employing particles coupled to a ds 100-mer oligonucleotide bearing a nick, a single-strand to 20 double-strand transition, a gap of 35 bp, or a 10 base insertion loop. Notably, under the assay conditions employed, ATM binds to these DNA molecules with equal efficiency and apparent affinity as it does to the fully ds DNA oligonucleotide (Figure 1C). Additional studies 25 show that ATM also binds effectively to ss DNA (Figure 1C) and that, as with ds-DNA, this binding is dependent on oligonucleotide length (data not shown). Furthermore. ATM binding in such experiments is competed effectively by linear and circular plasmid DNA, suggesting that DNA 30 termini are not required for ATM binding (NDL, unpublished data). Taken together, these data show that ATM, or a complex containing this factor, is capable of interacting with DNA molecules containing a variety of different structures in an apparently non-sequence 35 specific fashion. Our results also show that ATM prefers to bind to linear DNA, preferentially binding to the ends of the DNA.

Purification of ATM

To increase our understanding of ATM further, we decided 5 to attempt to purify this protein to essential homogeneity and thus separate it from other DNA-binding proteins, DNA repair factors, and protein and lipid The purification strategy we developed is outlined in Figure 2A. Since ATM is expressed ubiquitously and is located primarily in the cell 10 nucleus, HeLa cell nuclear extract was used as starting material. Because no biochemical assay was available for ATM protein function, we monitored its purification by Western blot analysis using antibodies raised against two different portions of the protein (Lakin et al., 1996). 15 This approach not only revealed the fractionation of ATM but also allowed us to pool fractions that were devoid of the abundant DNA-PK enzyme through simultaneously testing for the presence of DNA-PKcs and Ku. In light of the DNAbinding properties of ATM, we employed a final DNA 20 affinity step in the purification scheme (Figure 2B, lane Silver staining demonstrates that this leads to an essentially homogenous preparation of a ~350 kDa polypeptide, and Western blotting studies reveal that 25 this is recognised strongly by ATM antiserum ATM.B (Figure 2B). Since this protein is also recognised by two other antibodies raised to distinct regions of the ATM polypeptide (data not shown), we conclude that the purified protein is indeed ATM. As revealed in Figure 2B, whilst ATM is enriched throughout the purification 30 procedure, Ku, DNA-PK $_{cs}$, and the abundant ss DNA binding protein Replication Protein A (RPA) are all efficiently removed. Quantitative Western blotting and silverstaining reveal that the final yield of ATM is approximately 25% and indicate that ATM is of relatively 35 low abundance, comprising around 0.002% of total nuclear protein by weight.

Purified ATM possesses an associated p53 kinase activity Notably, as for DNA-PK (Hartley et al., 1995), purified ATM preparations were found to be devoid of detectable kinase activity towards PI and a variety of phosphorylated PI derivatives. Although we cannot 5 exclude that ATM phosphorylates these or related phospholipids under certain conditions or in the presence of additional components, we conclude that ATM is not a To assay for possible ATM-associated lipid kinase. protein kinase activity, we performed in vitro kinase 10 assays using equivalent amounts of various recombinant or purified proteins that we speculated may-be ATM substrates. Certain candidate substrates, such as DNA- PK_{cs} , Ku, proliferating cellular nuclear antigen (PCNA), and the 34 kDa subunit of RPA (RPA-p34), were chosen by 15 virtue of their association with DNA damage detection and/or involvement in DNA repair. We also tested Spl and p53, since these are both good substrates for DNA-PK and because A-T cells display aberrant induction of p53 in response to IR. A final protein tested was I,B, since 20 recent data have implicated this is an ATM target (Jung et al., 1995; Jung et al., 1997). Given that we had found that ATM binds to DNA, we included a DNA oligonucleotide known to activate DNA-PK in all initial 25 kinase reactions.

Notably, none of DNA-PK_{CS}, RPA-p45 and PCNA was phosphorylated efficiently by purified ATM (Figure 3A). However, longer exposures of autoradiograms reveals weak phosphorylation of both the 70 kDa subunit of Ku (Ku70) and Sp1 by ATM preparations (data not shown). Furthermore, prolonged exposures also reveal that ATM is capable of autophosphorylation (data not shown), consistent with previous rough studies employing ATM that had been immunoprecipitated directly from crude cell extracts (Keegan et al., 1996) (likely to contain all sorts of impurities). Most significantly, however,

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several independently purified ATM preparations were consistently found to phosphorylate p53 with high efficiency (Figure 3A) (contrary to the mentioned results of Keegan et al.). Taken together, these data reveal that, under our assay conditions, a protein kinase activity co-purifies with ATM that phosphorylates p53 efficiently, and Sp1 and Ku70 weakly. Importantly, DNA-PK efficiently phosphorylates p53, Sp1, Ku70 and RPA-p34 in vitro, revealing that the ATM-associated kinase activity exhibits a different substrate specificity from that of DNA-PK. This, together with the absence of detectable DNA-PK_{CS} or Ku in our ATM preparations argues strongly against the possibility that the ATM-associated protein kinase activity is imparted by DNA-PK contamination.

Although the above results reveal that a p53 kinase activity co-purifies with ATM, prolonged silver staining reveals additional polypeptides in our ATM preparations (data now shown). The possibility therefore existed that 20 the p53 kinase activity that we had detected was not mediated by ATM but by a contaminating protein. address this issue, we immunoprecipitated ATM from purified ATM preparations using polyclonal antibodies raised against either the N-terminal region (ATM.N) or an 25 internal region (ATM.B) of the ATM polypeptide (Lakin et al., 1996). After washing the immunoprecipitated material extensively in the presence of 500 mM KCl and 0.1% Nonidet-P40, it was employed in kinase reactions using p53 as substrate. To establish the purity of the 30 immunoprecipitated material, purified ATM was biotinylated and immunoprecipitated in parallel with ATM employed in the kinase reactions. The biotinylated precipitated proteins were then visualised by Western transfer and probing with streptavidin conjugated 35 horseradish peroxidase.

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As illustrated in Figure 3B, a biotinylated protein of approximately 350 kDa in size, the predicted molecular mass of ATM, is precipitated in these studies by anti-ATM antisera but not by pre-immune sera. Notably, no other proteins are consistently precipitated by both ATM antisera in these assays (a polypeptide of ~100 kDa is apparent in the ATM.N precipitation in Figure 3B but is not present in ATM.B immunoprecipitates and was not consistently observed in subsequent experiments using ATM.N).

Most importantly, these experiments revealed that p53 kinase activity is immunoprecipitated by the two ATM antisera. Greater ATM associated kinase activity is observed with ATM.N than with ATM.B, despite only slightly higher amounts of ATM being precipitated by ATM.N (Figure 3C). One possible explanation for this is that ATM.B, which recognises epitopes close to the ATM kinase domain, impairs ATM protein kinase activity. These studies show that the p53 kinase activity present in our ATM preparations follows ATM through a further

highly stringent immuno-affinity purification step, and suggest strongly that ATM directly mediates p53 phosphoylation. Although unlikely in our opinion, it remains a possibility that p53 is phosphorylated by a distinct polypeptide that has escaped our detection methods and which remains associated with ATM throughout the stringent purification and immunoprecipitation protocols employed.

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ATM associated kinase activity is stimulated by DNA Given that ATM can interact with DNA, we investigated whether ATM associated protein kinase activity is stimulated by a nucleic acid cofactor. To achieve this, we performed in vitro kinase assays using purified ATM either in the absence or presence of increasing amounts of DNA. Because previous studies have revealed that co-

localisation of DNA-PK and Sp1 to the same DNA molecule increases phosphorylation efficiency (Lees-Miller et al., .1992; Gottlieb and Jackson, 1993), we employed a linear plasmid molecule bearing multiple p53 binding sites. These studies revealed that DNA addition leads to marked 5 stimulation of p53 phosphorylation by DNA-PK (Figure 4A, Strikingly DNA addition was also found to result in marked stimulation of p53 phosphorylation in reactions containing ATM (Figure 4A, top). purified ATM preparations contain a DNA-stimulatable p53 10 kinase activity. Longer exposures of autoradiograms reveal that the ATM polypeptide is also subject to phosphorylation in such assays and that this phosphorylation is stimulated by DNA (data not shown). Experiments employing equimolar amounts of DNA-PK and ATM 15 revealed that the stimulation of p53 kinase activity by DNA is similar for ATM and DNA-PK, and that the stoichiometry of p53 phosphorylation by ATM is at least as high as that catalysed by DNA-PK (data not shown). Although DNA-dependent kinase activity was consistently 20 observed in ATM preparations, the degree of activation In this regard, additional polypeptides was variable. were apparent in several preparations that displayed high levels of DNA activatability. Thus, it is possible that 25 co-purifying polypeptides may be involved in high level ATM DNA dependent kinase activity. Notably, DNA-PK and ATM preparations both displayed significant but low levels of p53 kinase activity in the absence of DNA. is not currently known, however, whether this reflects bona fide DNA-independent phosphorylation or results from 30 small amounts of DNA in the protein preparations. Parallel experiments using cyclin A/cdk2 demonstrate no increase of p53 phosphorylation upon DNA addition (Figure 4A), and a variety of other protein kinases that we have tested are not stimulated by DNA. These results 35 therefore show that increased protein phosphorylation is not a general effect of adding DNA to p53 kinase assays

and reveal that ATM is highly unusual in its ability to be stimulated by DNA.

We had established that ATM binds to various types of linear DNA molecule (see Figure 1). Our binding 5 competition studies indicated that ATM also interacts with supercoiled and nicked DNA (data not shown). tested whether ATM associated kinase activity is affected differentially by various DNA structures. p53 kinase assays were performed in the absence of DNA or in the 10 presence of increasing amounts of either supercoiled or restriction enzyme-linearised plasmid DNA. Notably, ATM is activated by supercoiled and linear DNA (Figure 4B), and additional studies revealed that good activation also occurs with nicked plasmid DNA molecules 15 By contrast, DNA-PK is stimulated (data not shown). strongly by linear but only weakly by supercoiled plasmid DNA (Figure 4B; based on previous studies, the weak activation by the latter probably reflects small amounts 20 of nicked and/or linear DNA in the supercoiled plasmid preparation). These results are therefore consistent with data showing that ATM is able to interact with many different types of DNA structure. Furthermore, they show that, although ATM is analogous to DNA-PK in that its associated kinase activity is stimulated by DNA, the DNA 25 cofactor requirements of the two enzymes are different.

ATM associated kinase activity phosphorylates p53 at two sites

To determine the site(s) of p53 that are phosphorylated by ATM, bacterially expressed p53 was radioactively phosphorylated by ATM in either the presence or absence of DNA. Labelled p53 was purified by electrophoresis, digested by trypsin, and the resulting products separated by reverse-phase HPLC. Analysis of the resulting radioactive profiles showed a major peak eluting at 11-12% acetonitrile. A novel set of radioactive p53 derived

HPLC polypeptide peaks, which elute at 28-29% acetonitrile were induced substantially in the presence Phosphoamino acid analysis revealed that the DNA induced peaks contained peptides labelled at both serine and threonine residues, suggesting either two distinctly 5 labelled co-eluting peptides, or a single peptide containing both phosphoserine and phosphothreonine residues (data not shown). Radioactive peaks with similar elution properties were identified following phosphorylation of p53 by DNA-PK (Figure 5B) or casein 10 kinase I (data not shown). Previous studies have revealed that both DNA-PK and casein kinase I phosphorylate the N-terminal region of p53 (Lees-Miller et al., 1992; Milne et al., 1992). Initial attempts to sequence p53-derived peaks were unsuccessful, presumably 15 because they possess blocked amino-termini. However, cleavage with endoproteinase Asp-N allowed sequencing of Notably, release of counts at cycles 9 and 12 of Edman degradation of peptide 2a reveals that the sites of 20 phosphorylation correspond to p53 residues Ser-15 and Ser-15 has previously been demonstrated to be a phosphorylation site for DNA-PK (Lees-Miller et al., However, no detectable DNA-PK exists in our ATM preparations (see above).

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We therefore conclude that a novel DNA dependent kinase activity is associated with ATM that targets Ser-15 and Thr-18 of p53.

DNA-PK ATR has an associated kinase activity that phosphorylates p53 at Ser15 and Thr18

Given the fact than an activity in our ATM preparations was found to phosphorylate residue Thr18 of p53, we decided to test whether DNA-PK is also able to phosphorylate this site. To this end, p53 was incubated in the presence of radiolabelled [α³²P] ATP with purified human DNA-PK (a preparation consisting of the Ku and DNA-PK)

PKcs components of the enzyme; prepared as described in Hartley et al., 1995) in either the absence or presence of a linearised plasmid DNA molecule, then, as described for analysis of ATM-mediated phosphorylation events, the p53 was treated with protease to generate phospho-5 peptides and these were analysed by reverse-phase HPLC. These studies revealed that, as in the ATM studies, a set of related peptides eluting at around 28-29% acetonitrile (co-fractionating with ATM-derived peptides, 2a, b, and c; compare figures 5B and D) were phosphorylated by a 10 DNA-PK associated kinase activity in a DNA-inducible Furthermore, analysis of these revealed that they correspond to p53 peptides containing phosphorylation on residues Ser15 and Thr19 (Figure 5B). Subsequent studies using antibodies that recognise 15 specifically p53 that is phosphorylated on Ser15 or Thr19 (see below for details of antibody preparation) confirmed that the DNA-PK-associated kinase activity phosphorylates both of these residues of p53. Therefore, contrary to expectations, DNA-PK-associated kinase activity 20 phosphorylates p53 on Thr18 as well as Ser15.

ATR has an associated kinase activity that phosphorylates p53 at Ser15

Given that both DNA-PK-associated and a ATM-associated 25 kinase activities phosphorylate p53 on Ser15 and Thr18, we decided to see whether other kinases exist that can target these residues. To facilitate this approach, we generated rabbit polyclonal antibodies that specifically recognise p53 that is phosphorylated on Thr18 (they do 30 not recognise unphosphorylated p53 nor p53 that is phosphorylated solely on Ser15 nor is phosphorylated Similarly, we generated rabbit polyclonal elsewhere). antibodies that specifically recognised p53 phosphorylated on Serl5. These antibodies were generated 35 by immunising rabbits with specific p53-based phosphopeptides (containing either Thr18 or Ser15

phosphorylated), then preparing the antibodies with the desired recognition characteristics (those that recognised the specific phosphorylated peptides but not unphosphorylated versions of these peptides) by chromatography on columns bearing immobilised unphosphorylated peptide and columns bearing specific phosphorylated peptides.

To assess kinases activated in human cell extracts

capable of phosphorylating p53 on Ser15, HeLa nuclear
extract was fractionated chromatographically (see below)
then the resulting fractions were incubated with fulllength p53 protein and non-radioactively labelled ATP,
either in the absence or presence of DNA.

Phosphorylation of p53 was then assessed by subjecting
the samples to SDS-polyacrylamide cell electrophoresis are

the samples to SDS-polyacrylamide gel electrophoresis and Western immunoblotting. As shown in Figure 10, two main peaks of kinase activity (termed "activity 1" and "activity 2") capable to targeting Ser15 (S15) were detected in fractions of HeLa nuclear extract that had been chromatographed on Q-sepharose. Further analysis of these fractions revealed that both activities were

revealed that fractions comprising "activity 1" contained
the ATM-related protein ATR, whereas those comprising
"activity 2" contained DNA-PKcs (Figure 10). In
addition, other experiments revealed a third weaker,
activity peak in fractions between those comprising
activity 1 and activity 2, which corresponded to ATM.

stimulated by DNA. Furthermore, Western blotting

Further purification of activity peak 2 revealed that it corresponded to DNA-PK. Further fractionation of activity 1 revealed that, under all chromatographic separation techniques utilised, the DNA-activated p53 Ser15 kinase activity co-eluted with ATR. Indeed,

through following this kinase activity, ATR could be purified to near homogeneity (e.g. Figure 11; ATR was the only polypeptide whose elution was found to consistently

parallel that of the kinase activity). Thus, in addition to DNA-PK and ATM targeting p53 Ser15, we have made the surprising discovery that this residue is also phosphorylated by a kinase activity associated with ATR.

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Effect of p53 phosphorylation on interaction with Mdm2 To test whether phosphorylation on Ser15 or Thr18 of p53 affects its interaction with Mdm-2, phosphorylated and unphosphorylated p53-derived peptides were generated and were assessed for Mdm-2 binding by ELISA analysis. four peptides used contained p53 residues 11 to 25 (in the sequence NH2-SGSGEPPLSOETFSDLWKL-COOH; where the underlined sequence is that derived from p53) that were unphosphorylated (1); phosphorylated on residue equivalent to p53 residue Ser15(2); phosphorylated on residue equivalent to p53 residue Thr18(3); or phosphorylated on two residues, equivalent to p53 reside Ser15 and Thr18(4). Binding of Mdm-2 derivatives occurred effectively with unphosphorylated peptide 1 but was found to be inhibited dramatically in the cases of peptides 3 and 4, which contained phosphorylated Thr18. In contrast, binding was only impaired slightly by phosphorylation on Ser15 (peptide 2). We therefore conclude that phosphorylation on Thr18 of p53 has a dramatic effect on its interaction with Mdm-2 and that phosphorylation of this site is likely to play a key role in regulating p53 responses in vivo.

Additional purification method for ATM

HeLa nuclear extract was applied to Ni²⁺ - NTA agarose
(Qiagen). We found that ATM binds very tightly to this matrix, but not very well to Ni²⁺ - IDA matrices.

5 ml of nuclear extract was loaded onto a 1 x 2.5 cm column of Ni²⁺ - NTA agarose in the following buffer (Buffer D; 25 mM HEPES-KOH, pH 7.6, 100 mM KCl, 10% Glycerol, 1 mM MgCl₂, 20 mM imidazole). The column was

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washed extensively (10 column volumes) before applying a linear gradient of 20 mM - 500 mM imidazole in buffer D. Virtually pure ATM (as judged by silver stain analysis of 8% polyacrylamide gels) eluted near the end of the imidazole gradient. Less pure fractions of ATM eluted at the start of the gradient.

This provides a purification strategy for ATM or ATR that may be used alone, or in combination with various other chromatographic steps, e.g. DNA affinity chromatography as discussed already above.

Discussion

We have demonstrated that ATM is retained on immobilised particles bearing DNA molecules. Notably, ATM binds to 15 both ds and ss DNA in vitro, and studies employing a variety of unrelated oligonucleotides provide indication that this interaction is not sequence dependent. By exploiting these and other biochemical properties of ATM, we have developed a strategy to purify this polypeptide 20 from HeLa nuclear extracts to near homogeneity. The high purity of our final ATM preparations and the fact that ATM in such preparations can re-bind to DNA provides indication that ATM interacts with DNA directly. Although this appears somewhat different from the situation with 25 DNA-PKcs, which requires Ku to associate stably with DNA under our assay conditions, UV protein-DNA cross-linking has revealed that, in the context of the DNA-PKcs/Ku holoenzyme, DNA-PKcs does make close contacts with DNA (Gottlieb and Jackson, 1993). DNA-PKcs and ATM may 30 interact with DNA through similar mechanisms.

Because the C-terminal region of ATM possesses homology to the catalytic domain of mammalian PI 3-kinase, it has been speculated that ATM may phosphorylate inositol phospholipids. However, despite conducting lipid phosphorylation assays under various conditions and with

a variety of potential substrates, no ATM-associated lipid kinase activity was detected in our ATM preparations. These data are thus consistent with recent studies demonstrating that ATM-containing immunoprecipitates possess no detectable lipid kinase activity (Jung et al., 1997). Although we cannot discount the possibility that ATM modifies particular PI derivatives under certain conditions or in association with additional cofactors, we tentatively conclude that, as has been proposed for DNA-PKcs (Hartley et al., 1995) and FRAP (Brown et al., 1995), ATM is not a lipid kinase.

In contrast, our purified ATM preparations consistently possess protein serine/threonine kinase activity.

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Recently (Keegan et al., 1996) have performed rough experiments which might suggest that ATM-containing immunoprecipitates phosphorylate an ~350 kDa polypeptide, suggesting that ATM can modify itself (though the preparations would have contained all sorts of impurities, including kinases). We observe that purified ATM preparations are capable of some degree of ATM auto-phosphorylation.

In addition, we have tested ATM for its ability to modify 25 a variety of other polypeptides. Notably, despite the fact that IkB has been implicated as an ATM target by in vivo functional studies (Jung et al., 1995) and has recently been reported to be phosphorylated by ATM-containing immunoprecipitates (Jung et al., 1997), 30 under our assay conditions we do not detect significant IkB phosphorylation by ATM. Although alternatives exist, one explanation for this discrepancy is that IkB phosphorylation detected in the studies of (Jung et al., 1997) was mediated by a co-immunoprecipitating factor 35 that is separated from ATM during our purification scheme.

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Another protein that has been implicated as a possible ATM target by virtue of defective regulation in A-T cells is RPA (Liu and Weaver, 1993; Cheng et al., 1996). However, we have been unable to detect significant phosphorylation of RPA by ATM, suggesting that ATM regulates RPA indirectly. In contrast to the above, we observe low but detectable phosphorylation of Sp1 and the 70 kDa subunit of Ku by ATM. Although the significance of these phosphorylation events is uncertain, these findings raise the interesting possibilities that ATM plays a role in regulating Sp1-dependent transcription and controlling the activity of the Ku/DNA-PKcs holoenzyme.

By far the most efficient substrate for ATM that we have identified, however, is p53. Importantly, the p53 kinase activity we have detected consistently co-purifies with ATM, elutes from the final DNA affinity purification step with the same profile as the ATM polypeptide itself, and further co-purifies with ATM through an additional stringent immunoprecipitation procedure. These data provide strong indication that p53 kinase activity is an inherent property of the ATM polypeptide.

In a manner strikingly reminiscent of the activation of DNA-PK by DNA strand breaks and ds to ss DNA transitions, 25 we find that ATM and ATR associated p53 kinase activity is stimulated markedly by the addition of a DNA cofactor. There are several reasons why this DNA-stimulated protein kinase activity is unlikely to be mediated by contaminating DNA-PK. First, titration studies reveal 30 that, to provide the observed level of p53 phosphorylation, the DNA-PKcs content of ATM and ATR preparations would have to be essentially as great as that of ATM itself. Clearly, this is not the case silver staining and Western blotting reveal that, if any 35 residual DNA-PK does exist in our most purified ATM and ATR preparations, it is present at levels undetectable by

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the methods employed in this study. Second, the substrate specificity observed in ATM and ATR preparations is distinct from that of DNA-PK. Third, whereas ATM and ATR-associated kinase activity is stimulated similarly by supercoiled and linear plasmid molecules, DNA-PK is only activated strongly by the latter.

There are several possible ways in which ATM and ATR might be stimulated by DNA, and each of these may contribute to the effects that we observe. (The mechanism of action provides no limitation to the nature and scope of the present invention.)

One possibility is that DNA binding by ATM and ATR
activates the catalytic potential of the proteins
directly. Another is that the co-localisation of ATM and
ATR and its target DNA binding protein on the same DNA
molecule serves to potentiate interactions between the
kinase and its target. In line with one or both of the
above models, we have observed that ATM
auto-phosphorylation is also enhanced by DNA, albeit to a
lesser degree than that observed with p53.

Alternatively, at least part of the dramatic stimulation of p53 phosphorylation upon DNA addition could be explained by the binding of p53 to DNA inducing a conformational change in p53 that makes it a more effective ATM or ATR substrate. Thus, Ser-15 and Thr-18 might only become accessible to ATM after p53 is bound to DNA. In accordance with such a model, it is known that the conformation of p53 does change upon binding to DNA (Halazonetis et al., 1993), and it has been observed that several naturally occurring p53 mutants that are defective in sequence-specific DNA binding exhibit reduced phosphorylation at Ser-15 (Ullrich et al., 1993).

Given the DNA-PK paradigm, and because of the previously

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described role of ATM in DNA damage signalling, it might be tempting to speculate that ATM or ATR protein kinase activity in vivo is triggered by specific types of DNA damage or stalled DNA replication forks that occur in response to IR. However, unlike DNA-PK, which is activated strongly in vitro only by DNA molecules bearing perturbations in the DNA double-helix, we find that ATM interacts with all types of DNA structure that we have tested. It is, therefore, possible that ATM is active constitutively in mammalian cells. An alternative model, which we currently favour, is that ATM and ATR associate with other polypeptides rather like DNA-PKcs interacts with Ku, and it is the function of these additional components to restrict ATM or ATR activity under normal circumstances and only allow their activation after exposure to DNA damaging agents. In this regard, it is interesting to note that yeast genetic data indicate the S. cerevisiae and S. pombe homologues of ATM or ATR function in conjunction with other polypeptides in DNA damage signalling (reviewed in Elledge, 1996; Carr, 1997), and that biochemical studies reveal that ATM exists as a large complex of ~2 MDa in crude cell extracts (GCMS, unpublished data).

Together with genetic data indicating that ATM functions 25 upstream of p53 in a pathway for signalling IR-induced DNA damage, our findings provide indication that, following genomic insult, ATM and ATR phosphorylate p53 directly. Such a model would help to explain the deficient up-regulation of p53 in response to IR in A-T 30 cells and this, in turn, would explain at least some of the cell cycle checkpoint control defects of A-T cells. Interestingly, recent studies indicate that ATM interacts with p53 directly (Watters et al., 1997) providing a 35 possible mechanism for optimising the efficiency of ATM-mediated p53 phosphorylation in the cellular context. Indeed, since p53 itself binds to DNA strand breaks and

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DNA insertion loops (Balkalkin et al., 1994; Lee et al., 1995; Reed et al., 1995), p53 could actually play a role in targeting ATM or ATR to sites of DNA damage. model is particularly attractive when one considers that Ser-15 and Thr-18 reside in conserved and functionally important regions of the p53 polypeptide. Moreover, Ser-15 of p53 has been shown to be phosphorylated in vivo (reviewed in Anderson and Lees-Miller, 1992; Steegenga et al., 1996). In addition, although Thr-18 has not yet been identified as a physiological site for p53 modification, it is noteworthy that this residue is highly conserved in p53, and that around 8% of p53 phosphorylation in vivo occurs at Thr residues (Samad et al., 1986). In light of these points, it will clearly be of great interest to analyse the phosphorylation status of p53 Ser-15 and Thr-18 in wild-type and A-T cells, and to determine their degree of phosphorylation in response to IR.

Interestingly, phosphorylation of the N-terminal region 20 of p53 has been proposed to effect both the stability and the transcriptional activation potential of p53 (reviewed in Ko and Prives, 1996; Steegenga et al., 1996). Indeed, mutation of Ser-15 impairs the capacity of p53 to prevent S-phase progression and affects p53 stability (Fiscella 25 et al., 1993). Furthermore, p53 mutants unable to activate transcription show reduced phosphorylation at this site (Ullrich et al., 1993). Although no experiments have investigated the role of Thr-18 in p53 function directly, it is noteworthy that this residue forms part of the minimal p53 binding site for Mdm2, which functions 30 as a negative regulator of p53 function (Oliner et al., 1993). Significantly, Mdm-2 binding has been linked both to repressing p53-dependent transcriptional activation and targeting p53 for degradation within the cell (Momand et al., 1992; Oliner et al., 1993; Kubbutat et al., 35 1997). An attractive scenario, therefore, is that phosphorylation of p53 by ATM or ATR-may inhibit Mdm2

interaction, thus both stabilising p53 and de-repressing its transcriptional activity. Consistent with this, we find that the binding of p53-derived peptides to Mdm2 is strongly inhibited by phosphorylation of Thr18.

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It is emphasized that suggested mechanisms of action and models for ATM and p53 function discussed above are presented without limitation to the nature and scope of the present invention.

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EXPERIMENTAL PROCEDURES

DNA interaction studies

Oligonucleotides: one DNA strand containing a 5' biotin group (indicated by a "B" below) was annealed with 15 complementary oligonucleotide(s) and bound to streptavidin-coated iron-oxide particles (Dynabeads; Dynal, Oslo, Norway). HeLa nuclear extract, or ATM enriched extract (Q-Sepharose pool; see below) was incubated on ice for 30 min. with the DNA-iron oxide 20 particles. After washing with 5 x 0.5 ml of D* Buffer (25 mM HEPES-KOH, pH 7.6, 20% glycerol, 2 mM MgCl2, 0.2 mM EDTA, 1 mM DTT, 0.5 mM PMSF, 1mM Na Metabisulfite) containing 50 mM KC1, protein was eluted with 500 mM KC1 D* buffer or in gradual stepwise manner with KCl 25 concentrations of 100 mM, 250 mM and 500 mM in buffer D*. Fractions were analysed for ATM protein content by Western blotting using a previously described rabbit polyclonal antisera raised against amino acid residues 30 1980-2337 of ATM (Lakin et al., 1996).

Oligonucleotides:

ds 15-mer: 5' B-CCTGCCCTTGCCTGA-3'

5' TCAGGCAAGGGCAGG-3'

ds 25-mer 5' B-CCTGCCCTTGCCTGACGCTATTAGT-3'

5' ACTAATAGCGTCAGGCAAGGGCAGG-3'

ds 50-mer

- 5' B-TTGTAAAACGACGGCCAGTGAATTCATCATCAATAATATACCTTATTTTG-3'
- 5 CAAAATAAGGTATATTATTGATGATGAATTCACTGGCCGTCGTTTTACAA-3

'ds 75-mer

5 '5'

- 10 ss 50-mer
 - 5' B-TTGTAAAACGACGGCCAGTGAATTCATCATCAATAATATACCTTATTTTG-
- For the following, a biotinylated 100-mer oligonucleotide (DYNO) was used as a "backbone" to which other oligonucleotides were annealed.

DYNO 5' B-

CCTGCCCTTGCCTGACGCTATTAGTTCATCTATTTGTTTTGCTAATTCGA
TTGGAATCGAAACGGTCACATATTCTTTTTTTGACTGATTTCCTCGGCATA-3'

nicked oligo, DYNO + DAM2 + DAM3: ds/ss transition, DYNO + DAM3; gapped ds oligo, DYNO + DAM3 + DAM5; 10 bp insertion, DYNO + DAM6.

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DAM2:

5' TATGCCGAGGAAATCAGTCAAAAAAGAATATGTGACCGTTTCGATTCCAA-3'

DAM3:

30 5' TCGAATTAGCAAAACAAATAGATGAACTAATAGCGTCAGGCAAGGGCAGG-3'

DAM5: 5' TATGCCGAGGAAATC-3'

DAM6:

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TATGCCGAGGAAATCAGTCAAAAAAGAATATGTGACCGTTTCGAATTAGCAAAAC AAATAGATGAACTAATAGCGTCAGGCAAGGGCAGG-3'

ATM purification

All steps were performed at 4°C. HeLa nuclear extract .(20 ml) was applied to a Q-Sepharose column (35 ml, 1.5 \times .20 cm) equilibrated in D* buffer (25 mM HEPES-KOH, pH 7.6, 20% glycerol, 2 mM MgCl₂, 0.2 mM EDTA, 1 mM DTT, 0.5 5 mM PMSF, 1 mM Na Metabisulfite) containing 50 mM KCl. After washing with 2 column vol. of 50 mM KC1 D*, protein was eluted with a continuous salt gradient of 50 mM - 500 mM KC1 in D* buffer. ATM eluted between 160 and 200 mM 10 Fractions containing ATM and devoid of DNA-PK (as judged by Western blot analysis) were pooled and, after diluting to 100 mM KC1 in D* buffer, were loaded onto a heparin agarose column (1.5 \times 6 cm) pre-equilibrated in 100 mM KCl D* buffer. The column was washed with 2 column vol. of 100 mM KCl D* buffer before eluting with a 15 continuous gradient of 50 mM - 500 mM KCl in buffer D*. ATM was again followed by Western blot analysis and eluted between 200 and 220 mM KC1. Peak fractions were pooled and dialysed against 50 mM buffer D*. fractions were then incubated with gentle mixing for 1 h. 20 with 200 μ g biotinylated 50 bp ds DNA conjugated to streptavidin iron-oxide particles. Unbound protein was rebound to fresh DNA-iron oxide particles. were collected via a magnet and were washed 5 x with 0.5 ml of 50 mM KCl D* buffer before eluting ATM with 2 \times 75 25 μ l 500 mM KCl buffer D*. Purified ATM was snap-frozen and stored at -70°C.

ATR purification

ATR purification was carried out as set out in the description.

Immunological Methods

Western immunoblot analysis was performed as previously

(Lakin et al., 1996). Spl antisera were purchased from

Serotec Ltd. (Oxford, UK). RPA-p70 and RNA polymerase II

antisera were also utilised. Phospho-specific antisera

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were generated as described herein.

Immunoprecipitations were performed by incubating biotinylated or untreated purified ATM in parallel with serum for 1 h. on ice in D* buffer containing 50 mM KCl. Protein A Sepharose was added and the reaction incubated with slow rotation for a further h. at 4°C. Beads were washed at high stringency seven times in 500 μ l of D* buffer containing 500 mM KCl and 0.1% NP-40.

Biotinylated immunoprecipitated proteins were visualised by 7% SDS-PAGE followed by Western blotting and probing with streptavidin-conjugated horse-radish peroxidase. Un-biotinylated immunoprecipitated proteins were washed a further two times in 500 μ l 1 x Z' buffer prior to addition to kinase reactions (see below).

Phosphorylation assays

Kinase reactions were performed in 20 μ l containing: 10 μ l Z' buffer (25 mM HEPES-KOH pH 7.9, 50 mM KCl, 10 mM MgCl₂, 20% glycerol, 0.1% NP-40, 1mM DTT); 11 fmol ATM, DNA-PK or cyclin A/cdk2; 50-100 ng substrate and 0-30 fmol of DNA. Reactions were assembled and incubated for 3 min. on ice prior to addition of 10 μ Ci [γ -³²P] ATP and incubation at 30°C for 15 min. Phosphorylated proteins were subjected to 7% SDS-PAGE and visualised by autoradiography.

Mapping of p53 phosphorylation sites
Recombinant p53 (10-20 pmol; purified as previously (Hupp
30 et al., 1992)) was incubated with 12-24 ng of purified
ATM or DNA-PK in the presence of 100 μM ATP containing
10⁶-10⁷ cpm/nmol [³²P]-γATP under reaction conditions
described above. Linearised (pG₁₃-CAT) or supercoiled
(pBS-SK; Stratagene, USA) DNA were included in DNA-PK and
ATM reactions, respectively, where indicated. After 30
min. at 30°C, reactions were terminated by transferring
to an ice water bath. Following TCA-precipitation,

labelled p53 was resolved by 10% SDS-PAGE and visualised The gel section containing labelled by autoradiography. . p53 was excised and the protein eluted and TCA precipitated as described (Alessi et al., 1996). washed TCA pellet was either digested directly with 5 alkylated trypsin (Promega, Southampton, UK) or, for ASP-N digestion, solubilised first in 0.2% v/v Triton X-100 and digested overnight with 1:5 w:w Asp-N (Boehringer Mannheim) and, where indicated, followed by overnight digestion with trypsin. The supernatant containing digested protein was chromatographed on a Vydac 218TP54 C18 column (Separations Group, Hesperia, _CA) equilibrated with 0.1% v/v triflouroacetic acid (TFA), and eluted with a linear acetronitrile gradient. The flow rate was 0.8 ml/min. and 0.4 ml fractions were collected. Peak fractions were coupled covalently to a Sequelon acrylamide membrane and analysed on an Applied Biosystems 470A sequencer using the modified programme described by (Stokoe et al., 1992) to determine Edman degradation cycle numbers corresponding to radioactivity release.

Additional Purification of ATM
Purification using NTA has been described already above.

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Claims:

- 1. An assay method for a compound able to modulate the interaction between (i) ATM or a protein having an associated kinase activity and (ii) p53 or any protein having ATM phosphorylation sites homologous to those of p53, the method including the steps of:
- (a) bringing into contact a substance including a peptide fragment of (i) or a derivative, variant or analogue thereof, a substance including the relevant fragment of p53 or any protein having phosphorylation sites homologous to p53, or a variant, derivative or analogue thereof, and a test compound; and
- (b) determining interaction or binding between said substances and test compound.
- 2. An assay method for a compound able to modulate the interaction between (i) ATM or a protein having an associated kinase activity and (ii) p53 or any protein having ATM phosphorylation sites homologous to those of p53, the method including the steps of:
 - (a) bringing into contact a substance which includes at least a fragment of (i) which phosphorylates p53 or any protein having phosphorylation sites homologous to those of p53, a substance which includes at least a fragment of p53 or any protein, which includes a site which is phosphorylated by (i), and a test compound; and
 - (b) determining phosphorylation at said site.
- 3. An assay method for a compound able to affect p53 activity by modulation of interaction between (i) ATM or a protein having an associated kinase activity and (ii) p53, the method including the steps of:
- (a) bringing into contact a substance which is p53
 or a fragment, derivative, variant or analogue thereof, and a test compound; and
 - (b) determining p53 activity in the presence and

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absence of (i).

- 4. The method according to any one of claims 1 to 3 wherein the protein having an associated kinase activity is DNA-PK or ATR.
- 5. An agent capable of modulating interaction between

 (i) ATM or a protein having an associated kinase activity and (ii) p53 or any protein having ATM phosphorylation sites homologous to p53 obtained using a method according to any of claims 1 to 4.
- 6. An agent according to claim 5 capable of modulating ATM-mediated phosphorylation on p53 or said homologous sites.
 - 7. A peptide fragment of p53 capable of modulating interaction between ATM or a protein having an associated kinase activity and p53.
- 8. A peptide according to claim 7 capable of modulating phosphorylation of p53 by ATM or by a protein having an associated kinase activity.
- 9. A peptide according to claim 7 or claim 8 which has a sequence found in human p53 including Thr18.
 - 10. A peptide according to claim 7 or claim 8 which has a sequence found in human p53 including Ser15.
 - 11. A peptide according to any one of claims 7 to 10 wherein the protein having an associated kinase activity is DNA-PK or ATR.
- 12. A nucleic acid isolate encoding a peptide according to any of claims 7 to 11.

- 13. A peptide fragment of ATM or a protein having an associated kinase activity which is capable of modulating interaction between ATM or the protein and p53.
- 14. A peptide according to claim 13 capable of modulating phosphorylation of p53 by ATM or a protein having an associated kinase activity.
- 15. The peptide of claim 13 or claim 14 wherein the protein having an associated kinase activity is DNA-PK or ATR.
 - 16. A nucleic acid isolate encoding a peptide according to any one of claims 13 to 15.

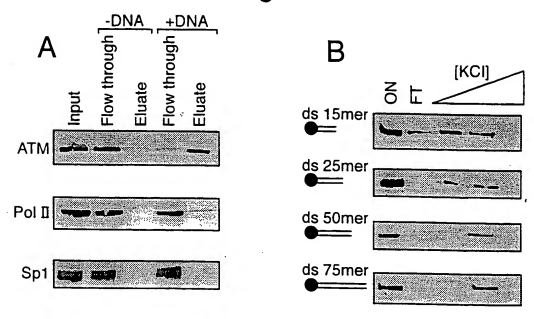
- 17. An agent or peptide fragment or nucleic acid isolate according to any of claims 5 to 16 for use in a method of treatment by therapy involving modulating ATM action.
- 18. Use of an agent or peptide fragment or nucleic acid isolate according to any of claims 5 to 17 in the manufacture of a medicament for modulating ATM action.
- 19. An assay method for a compound able to affect DNA binding by ATM or a protein having an associated kinase activity, the method including the steps of:
 - (a) bringing into contact a substance which is ATM or a protein having an associated kinase activity, or a fragment, variant or derivative which is able to bind DNA, DNA, and a test compound, under conditions wherein, in the absence of the test compound being an inhibitor of DNA binding by ATM or the protein having an associated kinase activity, said substance binds DNA; and
- (b) determining binding between said substance and 35 said DNA.
 - 20. The assay method according to claim 19 wherein the

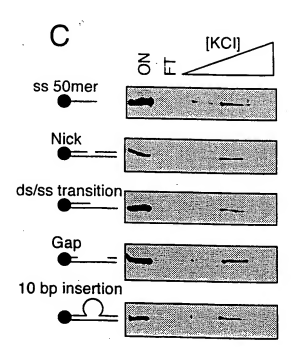
protein having an associated kinase activity is DNA-PK or ATR.

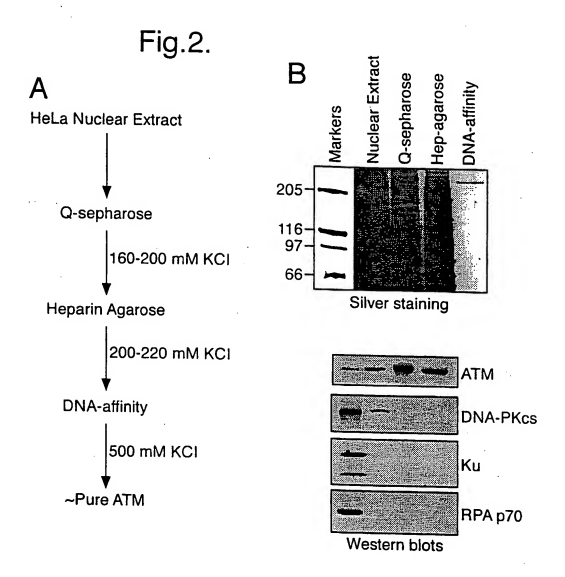
- 21. An agent capable of affecting DNA binding by ATMobtained using a method according to claim 19 or claim20.
 - 22. An agent according to claim 21 for use in a method of treatment by therapy involving modulating ATM action.
 - 23. Use of an agent according to claim 21 in the manufacture of a medicament for modulating ATM action.
- 24. A method of purifying ATM or ATR including
 contacting a mixture of molecules including ATM or ATR
 with DNA, washing off molecules which do not bind the
 DNA, and recovering ATM or ATR from the DNA-bound
 fraction.
- 20 25. Use of DNA for purifying ATM or ATR.
 - 26. A method of purifying ATM or ATR, including contacting a mixture of molecules including ATM or ATR with NTA, washing off molecules which do not bind the
- NTA, and recovering ATM or ATR from the NTA-bound fraction.
 - 27. A method according to claim 26 wherein the mixture is contacted with NTA in the presence of Ni^{2+} .
 - 28. Substantially pure ATM.
 - 29. Substantially pure ATR.

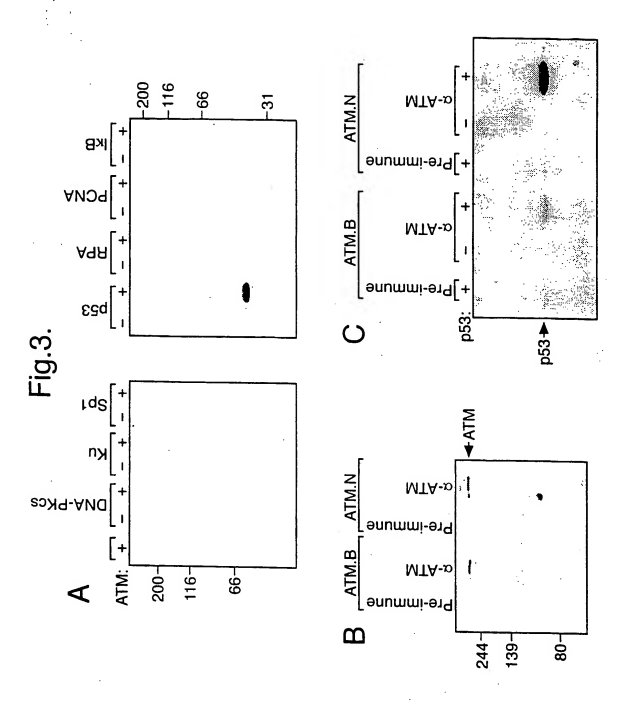
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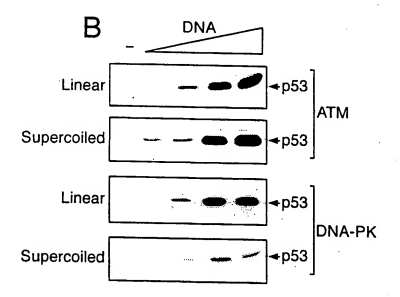
Fig.1.



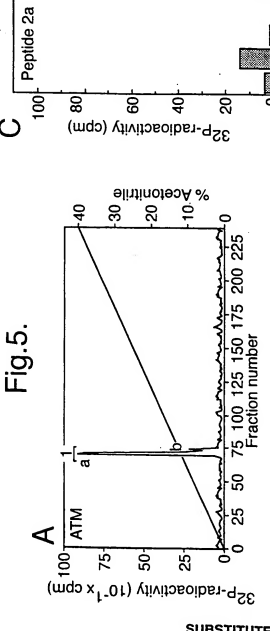


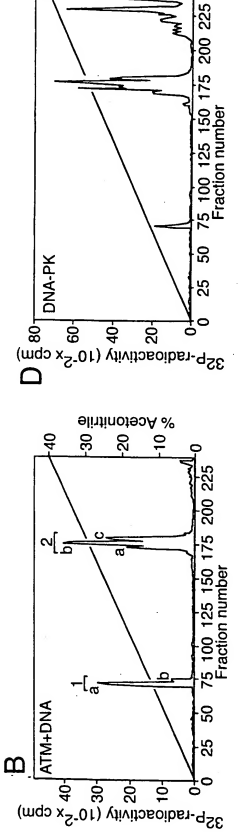


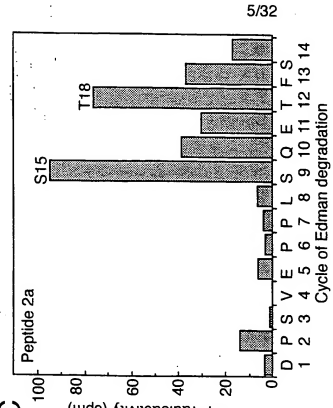




SUBSTITUTE SHEET (RULE 26)









% Acetonitrile

20

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KINASE DOMAIN

Figure 6b (1)

Sequence 9385 BP; 3030 A; 1685 C; 1973 G; 2697 T; 0 other;

Length: 9385 July 10, 1998 12:06 Type: N Check: 7765

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51 TTTGACCTIC CGAGTGCAGT GAGGCATACA TCACAATTTG GAATTATGCA

101 TTGGTTTATC AATTTACTTG TTTATTGTCA CCCTGCTGCC CAGATATGAC

TTCATGAGGA CAGTGATGTG TGTTCTGAAA TTGTGAACCA TGAGTCTAGT

1732
Figure 6b (11)
201 ACTTAATGAT CTGCTTATCT GCTGCCGTCA ACTAGAACAT GATAGAGCTA
251 CAGAACGAAA GAAAGAAGTT GAGAAATTTA AGCGCCTGAT TCGAGATCCT
301 GAAACAATTA AACATCTAGA TCGGCATTCA GATTCCAAAC AAGGAAAATA
351 TITGAATTGG GATGCTGTTT TTAGATTTTT ACAGAAATAT ATTCAGAAAG
401 AAACAGAATG TCTGAGAATA GCAAAACCAA ATGTATCAGC CTCAACACAA
451 GCCTCCAGGC AGAAAAAGAT GCAGGAAATC AGTAGTTTGG TCAAATACTT
501 CATCAAATGT GCAAACAGAA GAGCACCTAG GCTAAAATGT CAAGAACTCT
551 TAAATTATAT CATGGATACA GTGAAAGATT CATCTAATGG TGCTATTTAC
601 GGAGCTGATT GTAGCAACAT ACTACTCAAA GACATTCTTT CTGTGAGAAA
651 ATACTOGTGT GAAATATCTC AGCAACAGTG GTTAGAATTG TTCTCTGTGT
701 ACTICAGGCT CTATCIGAAA CCTICACAAG ATGITCATAG AGTITTAGIG
751 GCTAGAATAA TICATGCTGT TACCAAAGGA TGCTGTTCTC AGACTGACGG
801 ATTAAATTCC AAATTTTTGG ACTTTTTTTC CAAGGCTATT CAGTGTGCGA
851 GACAAGAAAA GAGCTCTTCA GGTCTAAATC ATATCTTAGC AGCTCTTACT
901 ATCITCCTCA AGACTITGGC TGTCAACTIT CGAATTCGAG TGTGTGAATT
951 AGGAGATGAA ATTCTTCCCA CTTTGCTTTA TATTTGGACT CAACATAGGC
1001 TTAATGATTC TTTAAAAGAA GTCATTATTG AATTATTTCA ACTGCAAATT
1051 TATATCCATC ATCCGAAAGG AGCCAAAACC CAAGAAAAAG GTGCTTATGA
1101 ATCAACAAAA TGGAGAAGTA TITTATACAA CITATATGAT CIGCTAGTGA
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1201 AATATTGCCG TCAAAGAAAA TITGATTGAA TTGATGGCAG ATATCTGTCA
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1451 CAAAGTATCC TGCAAGTTTA CCTAACTGTG AGCTGTCTCC ATTACTGATG
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1701 CTTTGGCTTA CTTGGAGCCA TAATICAGGG TAGTITAGTT GAGGITGACA
1751 GAGAATTCTG GAAGTTATTT ACTGGGTCAG CCTGCAGACC TTCATGTCCT
1801 GCAGTATGCT GTTTGACTTT GGCACTGACC ACCAGTATAG TTCCAGGAGC
1851 GGTAAAAATG GGAATAGAGC AAAATATGTG TGAAGTAAAT AGAAGCTTTT

Figure	6	b	(111)
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1901	CTTTAAAGGA	ATCAATAATC	AAATGGCTCT	TATTCTATCA	GTTAGAGGGT
1951					GTAATTTTCC
2001					AAAAACTGTA
2051					ACACCACCAA
2101					TATTTCTTCA
2151					
					GAATGTGGTA
2201			ATTGGCTTCT		
2251	GAATCACTGG	ATCGCTGTCT	TCTGGGATTA	TCAGAACAGC	TICIGAATAA
2301	TTACTCATCT	GAGATTACAA	ATTCAGAAAC	TCTTGTCCGG	TGTTCACGTC
2351	TTTTGGTGGG	TGTCCTTGGC	TGCTACTGTT	ACATGGGTGT	AATAGCTGAA.
2401	GAGGAAGCAT	ATAAGTCAGA	ATTATTCCAG	AAAGCCAACT	CTCTAATGCA
2451	ATCTCCACCA	GAAAGTATCA	CTCTGTTTAA	AAATAAGACA	AATGAGGAAT
2501	TCAGAATTGG	TTCCTTGAGA	AATATGATGC	AGCTATGTAC	ACGITGCITG
2551	AGCAACTGTA	CCAAGAAGAG	TCCAAATAAG	ATTGCATCTG	GCTTTTTCCT
2601	GCGATTGTTA	ACATCAAAGC	TAATGAATGA	CATTGCAGAT	ATTTGTAAAA
2651	GTTTAGCATC	CTTCATCAAA	AAGCCATTTG	ACCGTGGAGA	AGTAGAATCA
2701	ATGGAAGATG	ATACTAATGG	AAATCTAATG	GAGGTGGAGG	ATCAGTCATC
2751	CATGAATCTA	TTTAACGATT	ACCCTGATAG	TAGTGTTAGT	GATGCAAACG
2801	AACCTGGAGA	GAGCCAAAGT	ACCATAGGTG	CCATTAATCC	TTTAGCTGAA
2851	GAATATCTGT	CAAAGCAAGA	TCTACTTTTC	TTAGACATGC	TCAAGTTCTT
2901	GTGTTTGTGT	GTAACTACTG	CTCAGACCAA	TACTGTGTCC	TTTAGGGCAG
2951	CTGATATTCG	GAGGAAATTG	TTAATGTTAA	TTGATTCTAG	CACGCTAGAA
3001	CCTACCAAAT	CCCTCCACCT	GCATATGTAT	CTAATGCTTT	TAAAGGAGCT
3051	TCCTGGAGAA	GAGTACCCCT	TGCCAATGGA	AGATGTTCTT	GAACTTCTGA
3101	AACCACTATC	CAATGTGTGT	TCTTTGTATC	GTCGTGACCA	AGATGTTTGT
3151	AAAACTATTT	TAAACCATGT	CCTTCATGTA	GIGAAAAACC	TAGGTCAAAG
3201	CAATATGGAC	TCTGAGAACA	CAAGGGATGC	TCAAGGACAG	TITCTTACAG
3251	TAATTGGAGC	ATTTTGGCAT	CTAACAAAGG	AGAGGAAATA	TATATTCTCT
3301	GTAAGAATGG	CCCTAGTAAA	TTGCCTTAAA	ACTITICATE	AGGCTGATCC
3351	TTATTCAAAA	TGGGCCATTC	TTAATGTAAT	GGGAAAAGAC	TITCCTGTAA
3401	ATGAAGTATT	TACACAATTT	CTTGCTGACA	ATCATCACCA	AGTTCGCATG
3451	TTGGCTGCAG	AGTCAATCAA	TAGATTGTTC	CAGGACACGA	AGGGAGATTC
3501	TTCCAGGTTA	CTGAAAGCAC	TTCCTTTGAA	GCTTCAGCAA	ACAGCTTTTG

Figure 6b (iv)

; 355	1 AAAATGCATA CTTGAAAGCT CAGGAAGGAA TGAGAGAAAT GTCCCATAGT
. 360	L GCTGAGAACC CTGAAACTTT GGATGAAATT TATAATAGAA AATCTGTTTT
365	ACTGACGTTG ATAGCTGTGG TTTTATCCTG TAGCCCTATC TGCGAAAAAC
3701	
3751	
3801	
3851	
3901	
3951	•
4001	
4051	
4101	
4151	
4201	AGTAATTTAC CAGAGATTGT GGTGGAGTTA TTGATGACGT TACATGAGCC
4251	AGCAAATTCT AGTGCCAGTC AGAGCACTGA CCTCTGTGAC TTTTCAGGGG
4301	ATTIGGATCC TGCTCCTAAT CCACCTCATT TTCCATCGCA TGTGATTAAA
4351	GCAACATTTG CCTATATCAG CAATTGTCAT AAAACCAAGT TAAAAAGCAT
4401	TTTAGAAATT CTTTCCAAAA GCCCTGATTC CTATCAGAAA ATTCTTCTTG
4451	CCATATGTGA GCAAGCAGCT GAAACAAATA ATGTTTATAA GAAGCACAGA
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4551	AAGTGGCTTA GGAGGAGCTT GGGCCTTTGT TCTTCGAGAC GTTATTTATA
4601	CTTTGATTCA CTATATCAAC CAAAGGCCTT CTTGTATCAT GGATGTGTCA
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4701	AGCCGTGACT TACTGTAAGG ATGCTCTAGA AAACCATCTT CATGTTATTG
4751	TIGGTACACT TATACCCCTT GTGTATGAGC AGGTGGAGGT TCAGAAACAG
4801	GTATTOGACT TGTTGAAATA CTTAGTGATA GATAACAAGG ATAATGAAAA
	CCTCTATATC ACGATTAAGC TTTTAGATCC TTTTCCTGAC CATGTTGTTT
4901	TTAAGGATTT GCGTATTACT CAGCAAAAAA TCAAATACAG TAGAGGACCC
4951	TTTTCACTCT TGGAGGAAAT TAACCATTTT CTCTCAGTAA GTGTTTATGA
5001	TGCACTICCA TTGACAAGAC TTGAAGGACT AAAGGATCIT CGAAGACAAC
5051	TGGAACTACA TAAAGATCAG ATGGTGGACA TTATGAGAGC TTCTCAGGAT
	AATCCGCAAG ATGGGATTAT GGTGAAACTA GTTGTCAATT TGTTGCAGTT
	ATCCAAGATG GCAATAAACC ACACTGGTGA AAAAGAAGTT CTAGAGGCTG
5201	TTGGAAGCTG CTTGGGAGAA GTGGGTCCTA TAGATTTCTC TACCATAGCT

Figure 6b (v)

525	1 ATACAACATA GTAAAGATGC ATCTTATACC AAGGCCCTTA AGTTATTTGA
530	
535	
540	AAAAACATIT TAGCCACAAA GACTGGACAT AGTITCTGGG AGATITATAA
545	
5501	GAAAAAAGTT TITAGAAGTA CCCAGATTTG ACAAAGAAAA CCCTTTTGAA
5551	GGCCTGGATG ATATAAATCT GTGGATTCCT CTAAGTGAAA ATCATGACAT
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5651	GTGAAATTCT TCAATTATTA AAGCCAATGT GTGAAGTGAA
5701	TGTCAGACTG TACTTCCATA CTTGATTCAT GATATTTTAC TCCAAGATAC
5 7 51	AAATGAATCA TGGAGAAATC TGCTTTCTAC ACATGTTCAG GGATTTTTCA
5801	CCAGCTGTCT TCGACACTTC TCGCAAACGA GCCGATCCAC AACCCCTGCA
5851	CACACITATIC CGATGCTGTT TGGATAAAAA
5901	THE PROPERTY CIGITATION CTACATGAGA AGACAAAAGA
5951	TITAATGATG CITICIGGCT GGATTTAAAT
6001	ACCICAGICI IGIGCIGCIC ACTITACAGC
6051	TITACTCTAT GCAGAAATCT ATGCAGATAA GAAAAGTATG GATGATCAAG
6101	AGAAAAGAAG TCTTGCATTT GAAGAAGGAA GCCAGAGTAC AACTATTTCT
6151	AGCTTGAGTG AAAAAAGTAA AGAAGAAACT GGAATAAGTT TACAGGATCT
6201	TCTCTTAGAA ATCTACAGAA GTATAGGGGA GCCAGATAGT TTGTATGGCT
6251	GTGGTGGAGG GAAGATGTTA CAACCCATTA CTAGACTACG AACATATGAA
6301	CACGAAGCAA TGTGGGGCAA AGCCCTAGTA ACATATGACC TCGAAACAGC
6351	AATCCCCTCA TCAACACGCC AGGCAGGAAT CATTCAGGCC TTGCAGAATT
6401 6451	TGGGACTCTG CCATATTCTT TCCGTCTATT TAAAAGGATT GGATTATGAA
	AATAAAGACT GGTGTCCTGA ACTAGAAGAA CTTCATTACC AAGCAGCATG
6551	GAGGAATATG CAGTGGGACC ATTGCACTTC CGTCAGCAAA GAAGTAGAAG
6601	CONTRACTOR THETACARIG CICTACARIC TCTARGAGAC
6651	AGAGAATTCT CTACATTTTA TGAAAGTCTC AAATATGCCA GAGTAAAAGA
6701	ACTOGRAGAG ATGTGTAAGC GCAGCCTTGA GTCTGTGTAT TCGCTCTATC
6751	CCACACTTAG CAGGITGCAG GCCATTGGAG AGCTGGAAAG CATTGGGGAG
6801	CHOTCHGARGE TATATATAA
6851	CAGIGATITT AGTITICAGG
	AGCCTATCAT GGCTCTACGC ACAGTCATTT TOGAGATCCT GATGGAAAAG

Fig	ure 61	o (v1)				
	6901	GAAATGGACA	ACTCACAAAG	AGAATGTATT	AAGGACATTC	TCACCAAACA
	6951	CCTTGTAGAA	CTCTCTATAC	TGGCCAGAAC	TTTCAAGAAC	ACTCAGCTCC
	7001	CTGAAAGGGC	AATATITCAA	ATTAAACAGT	ACAATTCAGT	TAGCTGTGGA
	7051	GTCTCTGAGT	GGCAGCTGGA	AGAAGCACAA	GTATTCTGGG	CAAAAAAGGA
	7101	GCAGAGTCTT	GCCCTGAGTA	TTCTCAAGCA	AATGATCAAG	AAGTTGGATG
	7151	CCAGCTGTGC	AGCGAACAAT	CCCAGCCTAA	AACTTACATA	CACAGAATGT
	7201	CTGAGGGTTT	GTGGCAACTG	GTTAGCAGAA	ACGTGCTTAG	AAAATCCTGC
	7251	GGTCATCATG	CAGACCTATC	TAGAAAAGGC	AGTAGAAGTT	GCTGGAAATT
	7301	ATGATGGAGA	AAGTAGTGAT	GAGCTAAGAA	ATGGAAAAAT	GAAGGCATTT
	7351	CTCTCATTAG	CCCGGTTTTC	AGATACTCAA	TACCAAAGAA	TTGAAAACTA
	7401	CATGAAATCA	TCGGAATTTG	AAAACAAGCA	AGCTCTCCTG	AAAAGAGCCA
	7451	AAGAGGAAGT	AGGTCTCCTT	AGGGAACATA	AAATTCAGAC	AAACAGATAC
	7501	ACAGTAAAGG	TTCAGCGAGA	GCTGGAGTTG	GATGAATTAG	CCCIGCGIGC
	7551	ACTGAAAGAG	GATCGTAAAC	GCTTCTTATG	TAAAGCAGTT	GAAAATTATA
	7601	TCAACTGCTT	ATTAAGTGGA	GAAGAACATG	ATATGTGGGT	ATTCCGGCTT
	7651	TGTTCCCTCT	GGCTTGAAAA	TTCTGGAGTT	TCTGAAGTCA	ATGGCATGAT
	7701	GAAGAGAGAC	GGAATGAAGA	TTCCAACATA	TAAATTTTTG	CCTCTTATGT
	7751	ACCAATTGGC	TGCTAGAATG	GGGACCAAGA	TGATGGGAGG	CCTAGGATTT
	7801	CATGAAGTCC	TCAATAATCT	AATCTCTAGA	ATTTCAATGG	ATCACCCCCA
	7851	TCACACTTTG	TTTATTATAC	TGGCCTTAGC	AAATGCAAAC	AGAGATGAAT
	7901	TTCTGACTAA	ACCAGAGGTA	GCCAGAAGAA	GCAGAATAAC	TAAAAATGTG
	7951	CCTAAACAAA	GCTCTCAGCT	TGATGAGGAT	CGAACAGAGG	CTGCAAATAG
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	8101	TGGAAGACTC	AGAGAAAAGG	CATAAATATT	CCAGCAGACC	AGCCAATTAC
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	8351	ACCTGAGACA	AGATGCTGTC	ATGCAACAGG	TCTTCCAGAT	GTGTAATACA
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	8451	TTATAAGGTG	GTICCCCTCT	CTCAGCGAAG	TGGTGTTCTT	GAATGGTGCA
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	8551	CATAAAAGAT	ACAGGCCAAA	TGATTTCAGT	GCCTTTCAGT	GCCAAAAGAA

Figure 6b (vii)

8601	AATGATGGAG	GTGCAAAAAA	AGTCTTTTGA	AGAGAAATAT	GAAGTCTTCA
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8701	AAATTCTTGG	ATCCAGCTAT	TIGGTTIGAG	AAGCGATTGG	CTTATACGCG
8751	CAGTGTAGCT	ACTICTICTA	TIGTIGGITA	CATACTTGGA	CTTGGTGATA
8801	GACATGTACA	GAATATCTIG	ATAAATGAGC	AGTCAGCAGA	ACTIGTACAT
8851	ATAGATCTAG	GIGTIGCTTT	TGAACAGGGC	AAAATCCTTC	CTACTCCTGA
8901	GACAGTTCCT	TTTAGACTCA	CCAGAGATAT	TGTGGATGGC	ATGGGCATTA
8951	CGGGTGTTGA	AGGTGTCTTC	AGAAGATGCT	GTGAGAAAAC	CATGGAAGTG
9001	ATGAGAAACT	CTCAGGAAAC	TCTGTTAACC	ATTGTAGAGG	TCCTTCTATA
9051	TGATCCACTC	TTTGACTGGA	CCATGAATCC	TTTGAAAGCT	TTGTATTTAC
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9151	GACCAAGAAT	GCAAACGAAA	TCTCAGTGAT	ATTGACCAGA	GTTTCGACAA
9201	AGTAGCTGAA	CGTGTCTTAA	TGAGACTACA	AGAGAAACTG	AAAGGAGTGG
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9301	CAGGCCATAG	ACCCCAAAAA	TCTCAGCCGA	CITTTCCCAG	GATGGAAAGC
9351	TTGGGTGTGA	TCTTCAGTAT	ATGAATTACC	CTTTC	

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Figure 7a
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                                                                   Figure 7b (1)
    101 ACTGCCTTCC GGGTCACTGC CATGGAGGAG CCGCAGTCAG ATCCTAGCGT
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    201
    251 ATGCTGTCCC CGGACGATAT TGAACAATGG TTCACTGAAG ACCCAGGTCC
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Figure 7b (11)

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601	GGCCATCTAC AAGCAGTCAC AGCACATGAC GGAGGTTGTG AGGCGCTGCC
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Figure 8a (i)
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   DT
        20-MAY-1996 (Rel. 47, Last updated, Version 1)
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→ DE
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   KW
   XX
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        Karlene A. Cimprich, Chemistry, Harvard University, 12 Oxford Street,
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        Cambridge, MA 02138, USA
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  FŢ
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Figure 8b(1)

Figure 8a (ii)

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LERALHYLKNETEIELGSLLRQDFQGLHNELLLRIGEHYQQVFNGLSILASFASSDDPY OGPRDI ISPELMADYLOPKLLGILAFFNMOLLSSSVGIEDKKMALNSLMSLMKLMGPKH VSSVRVKMMTTLRTGLRFKDDFPELCCRAWDCFVRCLDHACLGSLLSHVIVALLPLIHI QPKETAAIFHYLIIENRDAVQDFLHEIYFLPDHPELKKIKAVLQEYRKETSESTDLQTT LQLSMKAIQHENVDVRIHALTSLKETLYKNQEKLIKYATDSETVEPIISQLVTVLLKGC QDANSQARLLCGECLGELGA I DPGRLDFSTTETOGKDFTFVTGVEDSSFAYGLLMELTR AYLAYADNSRAQDSAAYAIQELLSIYDCREMETNGPGHQLWRRFPEHVREILEPHLNTR YKSSQKSTDWSGVKKPIYLSKLGSNFAEWSASWAGYLITKVRHDLASKIFTCCSIMMKH DFKVTIYLLPHILVYVLLGCNQEDQQEVYAEIMAVLKHDDQHTINTQDIASDLCOLSTO TVFSMLDHLTQWARHKFQALKAEKCPHSKSNRNKVDSMVSTVDYEDYOSVTRFLDLIPO DTLAVASFRSKAYTRAVMHFESFITEKKQNIQEHLGFLQKLYAAMHEPDGVAGVSAIRK AEPSLKEQILEHESLGLLRDATACYDRAIOLEPDQIIHYHGVVKSMLGLGOLSTVITOV NGVHANRSEWTDELNTYRVEAAWKLSQWDLVENYLAADGKSTIWSVRLGQLLLSAKKRD ITAFYDSLKLVRAEQIVPLSAASFERGSYQRGYEYIVRLHMLCELEHSIKPLFQHSPGD SSOEDSLINWVARLEMTONSYRAKEPILALRRALLSLINKRPDYNEMVGECWLQSARVARK AGHHQTAYNALLNAGESRLAELYVERAKWLWSKGDVHQALIVLQKGVELCFPENETPPE GKNMLIHGRAMLLVGRFMEETANFESNAIMKKYKDVTACLPEWEDGHFYLAKYYDKLMP MVTDNKMEKQGDLIRYIVLHFGRSLQYGNQFIYQSMPRMLTLWLDYGTKAYEWEKAGRS DRVQMRNDLGKINKVITEHTNYLAPYQFLTAFSQLISRICHSHDEVFVVLMEIIAKVFL AYPQQAMWMTAVSKSSYPMRVNRCKEILNKAIHMKKSLEKFVGDATRLTDKLLELCNK PVDGSSSTLSMSTHFKMLKKLVEEATFSEILIPLOSVMIPTLPSILGTHANHASHEPFP GHWAYIAGFDDMVEILASLQKPKKISLKGSDGKFYIMMCKPKDDLRKDCRLMEFNSLIN KCLRKDAESRRRELHIRTYAVIPLNDECGIIEWVNNTAGLRPILTKLYKEKGVYMTGKE LRQCMLPKSAALSEKLKVFREFLLPRHPPIFHEWFLRTFPDPTSWYSSRSAYCRSTAVM SMVGYILGLGDRHGENILFDSLTGECVHVDFNCLFNKGETFEVPEIVPFRLTHNMVNGM GPMGTEGLFRRACEVIMRLMRDQREPLMSVLKTFLHDPLVEWSKPVKGHSKAPLNETGE VVNEKAKTHVLDIEQRLQGVIKTRNRVTGLPLSIEGHVHYLIQEATDENLLCOMYLGWT

Sequence 8210 BP; 2511 A; 1555 C; 1738 G; 2406 T; 0 other;

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- 301 GIGATGITGC TIGATITCAT CCAGCATATC ATGAAATCCT CCCCACTTAT

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- 401 TCAGTAATTG GATCATAACG AGACTTCTGC GGATTGCAGC AACTCCCTCC
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- 601 GGTCATGCTG TGGAATGGCC AGTGGTCATG AGCCGATTTT TAAGTCAATT
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Figure 8b (11)

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2251	TGTACTCTTC ACGGCATGIT TTATCTGACA AGITCTTTAA CAGAACCTTT
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Figure 8b (111)

2551	1 CCAGACAAAG ATGTTAGAGT GGCTTTT	'AGT GGAAATATCA AGCACATATT
2601	1 GGAATCCTTG GACTCTGAAG ATGGATT	TAT AAAGGAGCTT TTTGTCTTAA
2651	1 GAATGAAGGA AGCATATACA CATGCCC	AAA TATCAAGAAA TAATGAGCTG
2701	1 AAGGATACCT TGATTCTTAC AACAGGG	GAT ATTGGAAGGG CCGCAAAAGG
2751	1 AGATTIGGTA CCATTIGCAC TCTTACA	CTT ATTGCATTGT TTGTTATCCA
2801	1 AGTCAGCATC TGTCTCTGGA GCAGCAT	ACA CAGAAATTAG AGCTCTGGTT
2851	1 GCAGCTAAAA GTGTTAAACT GCAAAGT	TTT TTCAGCCAGT ATAAGAAACC
2901	CATCTGTCAG TTTTTGGTAG AATCCCT	TCA CTCTAGTCAG ATGACAGCAC
2951	TTCCGAATAC TCCATGCCAG AATGCTG	ACG TGCGAAAACA AGATGTGGCT
3001	CACCAGAGAG AAATGGCTTT AAATACG	ITG TCTGAAATTG CCAACGTTTT
3051	CGACTITCCT GATCITAATC GITTTCT	TAC TAGGACATTA CAAGTTCTAC
3101	TACCTGATCT TGCTGCCAAA GCAAGCC	CTG CAGCTTCTGC TCTCATTCGA
3151	ACTITAGGAA AACAATTAAA TGTCAAT	OGT AGAGAGATIT TAATAAACAA
3201	CTTCAAATAT ATTTTTTCTC ATTTGGT	TIG TICTIGITCC AAAGATGAAT
3251	TAGAACGTGC CCTTCATTAT CTGAAGAI	ATG AAACAGAAAT TGAACTGGGG
3301	AGCCTGTTGA GACAAGATTT CCAAGGAT	PTG CATAATGAAT TATTGCTGCG
3351	TATTGGAGAA CACTATCAAC AGGTTTT	TAA TGGTTTGTCA ATACTTGCCT
3401	CATTIGCATC CAGIGATGAT CCATATCA	AGG GCCCGAGAGA TATCATATCA
3451	CCTGAACTGA TGGCTGATTA TTTACAAC	CC AAATTGTTGG GCATTTTGGC
3501	TTTTTTAAC ATGCAGTTAC TGAGCTCT	AG TGTTGGCATT GAAGATAAGA
3551	AAATGGCCIT GAACAGITIG ATGTCITI	GA TGAAGTTAAT GGGACCCAAA
3601	and a contract of the contract	
3651	TCGATTCAAG GATGATTTTC CTGAATTC	
3701	TIGITOGCIG CCIGGATCAT GCITGICI	
3751	The state of the s	
	TATCTTCCAC TACCTCATAA TTGAAAAC	
3851	ccidal C	
	GCCGTTCTCC AGGAATACAG AAAGGAGA	
	The state of the s	
	The state of the s	
4151	AAGCTCGGTT GCTCTGTGGG GAATGTTT	AG GGGAATTGGG GGCGATAGAT

Figure 8b (iv)

4201	L CCAGGICGAT TAGATTICTC AACAACTGAA ACTCAAGGAA AAGATTITAC
4251	ATTTGTGACT GGAGTAGAAG ATTCAAGCTT TGCCTATGGA TTATTGATGG
4301	AGCTAACAAG AGCTTACCTT GCGTACGCTG ATAATAGCCG AGCTCAAGAT
4351	TCAGCTGCCT ATGCCATTCA GGAGTTGCTT TCTATTTATG ACTGTAGAGA
4401	GATGGAGACC AACGGCCCAG GTCACCAATT GTGGAGGAGA TTTCCTGAGC
4451	ATGTTCGGGA AATACTAGAA CCTCATCTAA ATACCAGATA CAAGAGTTCT
4501	CAGAAGTCAA CCGATTGGTC TGGAGTAAAG AAGCCAATTT ACTTAAGTAA
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4601	TTACAAAGGT TCGACATGAT CTTGCCAGTA AAATTTTCAC CTGCTGTAGC
4651	ATTATGATGA AGCATGATTT CAAAGTGACC ATCTATCTTC TTCCACATAT
4701	TCTGGTGTAT GTCTTACTGG GTTGTAATCA AGAAGATCAG CAGGAGGTTT
4751	ATGCAGAAAT TATGGCAGTT CTAAAGCATG ACGATCAGCA TACCATAAAT
4801	ACCCAAGACA TIGCATCIGA TCTGTGTCAA CICAGTACAC AGACTGTGTT
· 4851	CTCCATGCTT GACCATCTCA CACAGTGGGC AAGGCACAAA TTTCAGGCAC
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4951	TCAATGGTAT CTACTGTGGA TTATGAAGAC TATCAGAGTG TAACCCGTTT
5001	TCTAGACCTC ATACCCCAGG ATACTCTGGC AGTAGCTTCC TTTCGCTCCA
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5101	AAGCAAAATA TICAGGAACA TCTTGGATTT TTACAGAAAT TGTATGCTGC
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5401	TOGACAGATG AATTAAACAC GTACAGAGTG GAAGCAGCTT GGAAATTGTC
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	ATCACAGCTT TITATGACTC ACTGAAACTA GTGAGAGCAG AACAAATTGT
5601	ACCICITICA GCIGCAAGCT TIGAAAGAGG CTCCTACCAA CGAGGATAIG
5701	AATATATTGT GAGATTGCAC ATGTTATGTG AGTTGGAGCA TAGCATCAAA
	CCACTITICC AGCATICICC AGGIGACAGI TCTCAAGAAG ATTCTCTAAA
5801	CTGGGTAGCT CGACTAGAAA TGACCCAGAA TTCCTACAGA GCCAAGGAGC
	CTATOCIGGC TCTCCCGAGG GCTTTACTAA GCCTCAACAA AAGACCAGAT TACAATGAAA TGGTTGGAGA ATGCTGGCIG CAGAGTGCCA GGGTAGCTAG

Figure 8b (v)

5901	AAAGGCTGGT	CACCACCAGA	CAGCCTACAA	TGCTCTCCTT	AATGCAGGGG
5951	AATCACGACT	CGCTGAACTG	TACGTGGAAA	GGGCAAAGTG	GCTCTGGTCC
6001	AAGGGTGATG	TTCACCAGGC	ACTAATIGIT	CTTCAAAAAG	GTGTTGAATT
6051	ATGITTICCT	GAAAATGAAA	CCCCACCTGA	GGGTAAGAAC	ATGTTAATÇC
6101	ATGGTCGAGC	TATGCTACTA	GTGGGCCGAT	TTATGGAAGA	AACAGCTAAC
6151	TTTGAAAGCA	ATGCAATTAT	GAAAAAATAT	AAGGATGTGA	CCGCGTGCCT
6201	GCCAGAATGG	GAGGATGGGC	ATTTTTACCT	TGCCAAGTAC	TATGACAAAT
6251	TGATGCCCAT	GGTCACAGAC	AACAAAATGG	AAAAGCAAGG	TGATCTCATC
6301	CGGTATATAG	TICTTCATTT	TGGCAGATCT	СТАСААТАТС	GAAATCAGTT
6351	CATATATCAG	TCAATGCCAC	GAATGTTAAC	TCTATGGCTT	GATTATGGTA
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6451	AGGAATGATT	TGGGTAAAAT	AAACAAGGTT	ATCACAGAGC	АТАСАААСТА
6501	TTTAGCTCCA	TATCAATTIT	TGACTGCTTT	TTCACAATTG	ATCTCTCGAA
6551	TITGTCATTC	TCACGATGAA	GTTTTTGTTG	TCTTGATGGA	AATAATAGCC
6601	AAAGTATTTC	TAGCCTATCC	TCAACAAGCA	ATGTGGATGA	TGACAGCTGT
6651	GTCAAAGTCA	TCTTATCCCA	TGCGTGTGAA	CAGATGCAAG	GAAATCCTCA
6701	ATAAAGCTAT	TCATATGAAA	AAATCCTTAG	AGAAGTTTGT	TGGAGATGCA
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6851	TGGTAGAAGA	AGCAACATTT	AGTGAAATCC	TCATTCCTCT	ACAATCAGTC
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6951	CCATGAACCA	TTTCCTGGAC	ATTGGGCCTA	TATTGCAGGG	TTTGATGATA
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7151			CGTAGAAGAG		
7201			TGAATGTGGG		
7251			TTCTGACCAA		
7301			CTTCGCCAGT		
7351			AGTATTCCGA		
7401			GGTTTCTGAG		
7451			GCTTACTGCC		
7501	ATGGTTGGTT	ATATICIGGG	GCTTGGAGAC	CCTCATCCTC	AAAATATTCT

Figure 8b (v1)

7551	CTTTGATTCT	TTGACTGGTG	AATGCGTACA	TGTAGATTTC	AATTGTCTTT
7601	TCAATAAGGG	AGAAACCTTT	GAAGTTCCAG	AAATTGTGCC	ATTICGCCTG
7651	ACTCATAATA	TGGTTAATGG	AATGGGTCCT	ATGGGAACAG	AGGGTCTTTT
7701	TCGAAGAGCA	TGTGAAGTTA	CAATGAGGCT	GATGCGTGAT	CAGCGAGAGC
7751	CTTTAATGAG	TGTCTTAAAG	ACTITICTAC	ATGATCCTCT	TGTGGAATGG
7801	AGTAAACCAG	TGAAAGGGCA	TTCCAAAGCG	CCACTGAATG	AAACTGGAGA
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8101	GTGGTTGTAT	CTGTTCAATT	CTAAAGTACA	ACATAAATTT	ACGTTCTCAG
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8201	CACTCGTGCC				

Figure 9a (1)

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22-FEB-1997 (Rel. 51, Last updated, Version 4)
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       *DWA-dependent protein kinase catalytic subunit: a relative of
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Figure 9 (11)

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FT KWKVPTYKDYVDLFRHLLSSDQMMDSILADEAFFSVNSSSESLNHLLYDEFVKSVLKIV EKLDLTLEIQTVGEQENGDEAPGVWMIPTSDPAANLHPAKPKDFSAFINLVEFCREILP FT FT EKQAEFFEPWVYSFSYELILQSTRLPLISGFYKLLSITVRNAKKIKYFEGVSPKSLKHS PEDPEKYSCFALFVKFGKEVAVKMKQYKDELLASCLTFLLSLPHNIIELDVRAYVPALQ FT MAFKLGLSYTPLAEVGLNALEEWSIYIDRHVMQPYYKDILPCLDGYLKTSALSDETKNN FT WEVSALSRAAQKGFNKVVLKHLKKTKNLSSNEAISLEEIRIRVVQMLGSLGGQINKNLL FT FT TVTSSDEMMKSYVAWDREKRLSFAVPFREMKPVIFLDVFLPRVTELALTASDROTKVAA CELLHSMVMFMLGKATQMPEGGQGAPPMYQLYKRTFPVLLRLACDVDQVTRQLYEPLVM FT FT QLIHWFTNNKKFESQDTVSLLEAILDGIVDPVDSTLRDFCGRCIREFLKWSIKQITPQQ QEKSPVNTKSLFKRLYSLALHPNAFKRLGASLAFNNIYREFREEESLVEQFVFEALVIY FT FT MESLALAHADEKSLGTIQQCCDAIDHLCRIIEKKHVSLNKAKKRRLPRGFPPSASLCLL DLVKWLLAHCGRPQTECRHKSIELFYKFVPLLPGNRSPNLWLKDVLKEEGVSFLINTFE FT GGGCGQPSGILAQPTLLYLRGPFSLQATLCWLDLLLAALECYNTFIGERTVGALQVLGT FT EAQSSLLKAVAFFLESIAMHDIIAAEKCFGTGAAGNRTSPQEGERYNYSKCTVVVRIME FT FTTTLLNTSPEGWKLLKKDLCNTHLMRVLVQTLCEPASIGFNIGDVQVMAHLPDVCVNL FT MKALKMSPYKDILETHLREKITAQSIEELCAVNLYGPDAQVDRSRLAAVVSACKQLHRA FT GLLHNILPSQSTDLHHSVGTELLSLVYKGIAPGDERQCLPSLDLSCKQLASGLLELAFA FT FGGLCERLVSLLLNPAVLSTASLGSSQGSVIHFSHGEYFYSLFSETINTELLKNLDLAV FT LELMQSSVDNTKMVSAVLNGMLDQSFRERANQKHQGLKLATTILQHWKKCDSWWAKDSP FT LETKMAVLALLAKILQIDSSVSFNTSHGSFPEVFTTYISLLADTKLDLHLKGQAVTLLP FT FFTSLTGGSLEELRRVLEQLIVAHFPMQSREFPPGTPRFNNYVDCMKKFLDALELSQSP FT MLLELMTEVLCREQQHVMEELFQSSFRRIARRGSCVTQVGLLESVYEMFRKDDPRLSFT FT RQSFVDRSLLTLLWHCSLDALREFFSTIVVDAIDVLKSRFTKLNESTFDTQITKKMGYY FΤ KILDVMYSRLPKDDVHAKESKINQVFHGSCITEGNELTKTLIKLCYDAFTENMAGENQL FT LERRRLYHCAAYNCAISVICCVFNELKFYQGFLFSEKPEKNLLIFENLIDLKRRYNFPV FT EVEVPMERKKKYIEIRKEAREAANGDSDGPSYMSSLSYLADSTLSEFMSQFDFSTGVQS FT YSYSSQDPRPATGRFRRREQRDPTVHDDVLELEMDELNRHECMAPLTALVKHMHRSLGP FT POGEEDSVPRDLPSWMKFLHGKLCNPIVPLNIRLFLAKLVINTEEVFRPYAKHWLSPLL FT FT QLAASENNGGEGIHYMVVEIVATILSWIGLATPIGVPKDEVLANRLLNFLMKHVFHPKR AVFRHNLEIIKTLVECWKDCLSIPYRLIFEKFSGKDPNSKDNSVGIQLLGIVMANDLPP FT FT YDPQCGIQSSEYFQALVNNMSFVRYKEVYAAAAEVLGLILRYVMERKNILEESLCELVA FT KQLKQHQNTMEDKFIVCLNKVTKSFPPLADRFMNAVFFLLPKFHGVLKTLCLEVVLCRV DGMTELYFQLKSKDFVQVMRHRDERQKVCLDIIYKMMPKLKPVELRELLNPVVEFVSHP FT FT STICREOMYNILMWIHDNYRDPESETDNDSQEIFKLAKDVLIQGLIDENPGLQLIIRNF FT WSHETRLPSNTLDRLLALNSLYSPKIEVHFLSLATNFLLEMTSMSPDYPNPMFEHPLSE FT CEFQEYTIDSDWRFRSTVLTPMFVETQASQGTLQTRTQEGSLSARWPVAGQIRATQQQH DFTLTQTADGRSSFDWLTGSSTDPLVDHTSPSSDSLLFAHKRSERLQRAPLKSVGPDFG FT FT KKRLGLPGDEVDNKVKGAAGRTDLLRLRRRFMRDQEKLSLMYARKGVADQKREKEIKSE FT LKMKQDAQVVLYRSYRHGDLPDIQIKHSSLITPLQAVAQRDPIIAKQLFSSLFSGILKE MDKFKTLSEKNNITQKLLQDFNRFLNTTFSFFPPFVSCIQDISCQHAALLSLDPAAVSA FT FT GCLASIQQPVGIRLLEFALIRLLPAELPAKRVRGKARLPPDVLRWVELAKLYRSIGEYD FT VLRGIFTSEIGTKQITQSALLAEARSDYSEAAKQYDEALNKQDWVDGEPTEAEKDFWEL ASLICYNHLAEWKSLEYCSTASIDSENPPDLNKIWSEPFYQETYLPYMIRSKLKLLLQG FT EADQSLLTFIDKAMHGELQKAILELHYSQELSLLYLLQDDVDRAKYYIQNGIQSFMQNY FT FT SSIDVLLHQSRLTKLQSVQALTEIQEFISFISKQGNLSSQVPLKRLLNTWINRYPDAKM DPMNIWDDIITNRCFFLSKIEEKLTPLPEDNSMNVDQDGDPSDRMEVQEQEEDISSLIR FT FT SCKFSMKMKMIDSARKQNNFSLAMKLLKELHKESKTRDDWLVSWVQSYCRLSHCRSRSQ FT GCSEQVLTVLKTVSILDEMNVSSYLXKNILAFRDONILLGTTYRIIANALSSEPACLAE IEEDKARRILELSGSSSEDSEKVIAGLYQRAFQHLSEAVQAAEEEAQPPSWSCGPAAGV FT IDAYMTLADFCDQQLRKEEENASVIDSAELQAYPALVVEKMLKALKLNSNEARLKFPRL FT FT LQIIFRYPEETLSLMTKEISSVPCWQFISWISHMVALLDKDQAVAVQHSVEEITDNYPQ FT AIVYPFIISSESYSFKDTSTGHKNKEFVARIKSKLDQGGVIQDFINALDQLSNPELLFK DWSNDVRAELAKTPVNKKNIEKMYERMYAALGDPKAPGLGAFRRKFIQTFGKEFDKHPG FT KGGSKILLRMKLSDFNDITNMLLLKMNKDSKPPGNLKECSPWMSDFKVEFLRNELEIPGQ FT FT YDGRGKPLPEYHVRIAGFDERVIVMASLRRPKRIIIRGHDEREHPFLVKGGEDLRODOR FT VEQLFQVMNGILAQDSACSQRALQLRTYSVVPMTSSDPRAPPCEYKLWLTKMSGKHDVG AYMLMYKGANRTETVTSFRKRESKVPADLLKRAFVRMSTSPEAFLALRSHFASSHALIC FT FT ISHWILGIGDRHLNNFMVAMETGGVIGIDFGHAFGSATQFLPVPELMPFRLTRQFINLM LPMKETGLMYS IMVHALRAFRSDPGLLTNTMDVFVKEPSFDWKNFEQKMLKKGGSWIQE FT INVAEKNWYPROKICYAKRKLAGANPAVITCDELLLGHEKAPAFRDYVAVARGSKDHNI FT FT RAQEPESGLSEETQVKCLMDQATDPNILGRTWEGWEPWM* XX Sequence 12780 BP; 3612 A; 2769 C; 3084 G; 3314 T; 1 other;

U34994 Length: 12780 July 10, 1998 12:15 Type: N Check: 8189 .. Figure 9b(1)

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Figure 9b (11)

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201	TGCTGGCATT	ACAGACATCT	TTAGTTTTTT	CCAGAGATTT	CGGTTTGCTT
251	GTATTTGTCC	GGAAGTCACT	CAACAGTATT	GAATTTCGTG	AATGTAGAGA
301	AGAAATCCTA	AAGTTTTTAT	GTATTTTCTT	AGAAAAAATG	GGCCAGAAGA
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901	CACCTGCCTT	CTGGACAACT	ACGIGICICI	ATTTGAAGTC	TTGTTAAAGT
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1551	GCCCTGAGTC	TGAATCTGAA	GACCACCGTG	CTTCAGGGGA	AGTCAGAACT
1601	GGCAAATGGA	AGGTGCCCAC	ATACAAAGAC	TACGTGGATC	TCTTCAGACA
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25/32

Figure 9b (111)

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2551	TCTGAAGAAG ACAAAGAACC TTTCATCAAA CGAAGCAATA TCCTTAGAAG
2601	AAATAAGAAT TAGAGTAGTA CAAATGCTTG GATCTCTAGG AGGACAAATA
2651	AACAAAAATC TICIGACAGT CACGICCICA GATGAGATGA TGAAGAGCIA
2701	TGTGGCCTGG GACAGAGAGA AGCGGCTGAG CTTTGCAGTG CCCTTTAGAG
2751	AGATGAAACC TGTCATTTTC CTGGATGTGT TCCTGCCTCG AGTCACAGAA
2801	TTAGCGCTCA CAGCCAGTGA CAGACAAACT AAAGTTGCAG CCTGTGAACT
2851	TITIACATAGC ATGGTTATGT TTATGTTGGG CAAAGCCACG CAGATGCCAG
2901	AAGGGGGACA GGGAGCCCCA CCCATGIACC AGCICTATAA GCGGACGITT
	CCTGTGCTGC TTCGACTTGC GTGTGATGTT GATCAGGTGA CAAGGCAACT
3001	CINGTIANGE AGENCATICA CIGGITCACT AACAACAAGA
	AATTTGAAAG TCAGGATACT GTTTCCTTAC TAGAAGCTAT ATTGGATGGA
3101	TACTITIAGA GATTITIGIG GICGGIGIAT
3151	CCATTAAGCA AATAACACCA CAGCAGCAGG
3201	TOTAL
	GCGCTTCACC CCAATGCTTT CAAGAGGCTG GGAGCATCAC TTGCCTTTAA
	TAATATCTAC AGGGAATTCA GGGAAGAAGA GTCTCTGGTG GAACAGTTTG
	TGTTTGAAGC CTTGGTGATA TACATGGAGA GTCTGGCCTT- AGCACATGCA

Figure 9b (1v)

3401	GATGAGAAGT	CCTTAGGTAC	AATTCAACAG	TGTTGTGATG	CCATTGATCA
3451	CCTATGCCGC	ATCATTGAAA	AGAAGCATGT	TTCTTTAAAT	AAAGCAAAGA
3501	AACGACGTTT	GCCGCGAGGA	TTTCCACCTT	CCGCATCATT	GTGTTTATTG
3551	GATCTGGTCA	AGTGGCTTTT	AGCTCATTGT	GGGAGGCCCC	AGACAGAATG
3601	TCGACACAAA	TCCATTGAAC	TCTTTTATAA	ATTCGTTCCT	TTATTGCCAG
3651	GCAACAGATC	CCCTAATTTG	TGGCTGAAAG	ATGTTCTCAA	GGAAGAAGGT
3701	GTCTCTTTTC	TCATCAACAC	CTTTGAGGGG	GGTGGCTGTG	GCCAGCCCTC
3751	GGGCATCCTG	GCCCAGCCCA	CCCTCTTGTA	CCTTCGGGGG	CCATTCAGCC
3801	TGCAGGCCAC	GCTATGCTGG	CTGGACCTGC	TCCTGGCCGC	GTTGGAGTGC
3851	TACAACACGT	TCATTGGCGA	GAGAACTGTA	GGAGCGCTCC	AGGTCCTAGG
3901	TACTGAAGCC	CAGTCTTCAC	TTTTGAAAGC	AGTGGCTTTC	TTCTTAGAAA
3951	GCATTGCCAT	GCATGACATT	ATAGCAGCAG	AAAAGTGCTT	TGGCACTGGG
4001	GCAGCAGGTA	ACAGÁACAAG	CCCACAAGAG	GGAGAAAGGT	ACAACTACAG
4051	CAAATGCACC	GITGTGGTCC	GGATTATGGA	GTTTACCACG	ACTCTGCTAA
4101	ACACCTCCCC	GGAAGGATGG	AAGCTCCTGA	AGAAGGACTT	GTGTAATACA
4151	CACCTGATGA	GAGTCCTGGT	GCAGACGCTG	TGTGAGCCCG	CAAGCATAGG
4201	TTTCAACATC	GGAGACGTCC	AGGTTATGGC	TCATCTTCCT	GATGTTTGTG
4251	TGAATCTGAT	GAAAGCTCTA	AAGATGTCCC	CATACAAAGA	TATCCTAGAG
4301	ACCCATCTGA	GAGAGAAAAT	AACAGCACAG	AGCATTGAGG	AGCTTTGTGC
4351	CGICAACITG	TATGGCCCTG	ACGCGCAAGT	GGACAGGAGC	AGGCTGGCTG
4401	CIGITGIGIC	TGCCTGTAAA	CAGCTTCACA	GAGCTGGGCT	TCTGCATAAT
4451	ATATTACCGT	CTCAGTCCAC	AGATTTGCAT	CATTCTGTTG	GCACAGAACT
4501	TCTTTCCCTG	GTTTATAAAG	GCATTGCCCC	TGGAGATGAG	AGACAGIGIC
4551		AGACCTCAGT			
4601		CTTTTGGAGG			
4651		GTGCTGTCCA			
4701		CTCCCATGGG			
4751		AATTATTGAA			
4801		GTGGATAATA			•
4851		GAGCTTCAGG			
4901		CTACAATTCT			
4951		TCCCCTCTCG	•		
5001	CAAAAATTTT	ACAGATTGAT	TCATCIGTAT	CTTTTAATAC	AAGTCATOGT

Figure 9b (v)

5051	TCATTCCCTG	AAGTCTTTAC	AACATATATT	AGTCTACTTG	CTGACACAAA
5101	GCTGGATCTA	CATTTAAAGG	GCCAAGCTGT	CACTCTTCTT	CCATTCTTCA
5151	CCAGCCTCAC	TGGAGGCAGT	CTGGAGGAAC	TTAGACGTGT	TCTGGAGCAG
5201	CTCATCGTTG	CTCACTTCCC	CATGCAGTCC	AGGGAATTTC	CTCCAGGAAC
5251	TCCGCGGTTC	AATAATTATG	TGGACTGCAT	GAAAAAGTTT	CTAGATGCAT
5301	TGGAATTATC	TCAAAGCCCT	ATGTTGTTGG	AATTGATGAC	AGAAGTTCTT
5351	TGTCGGGAAC	AGCAGCATGT	CATGGAAGAA	TTATTTCAAT	CCAGTTTCAG
5401	GAGGATTGCC	AGAAGGGGTT	CATGTGTCAC	ACAAGTAGGC	CTTCTGGAAA
5451	GCGTGTATGA	AATGTTCAGG	AAGGATGACC	CCCGCCTAAG	TTTCACACGC
5501	CAGTCCTTTG	TGGACCGCTC	CCTCCTCACT	CTGCTGTGGC	ACTGTAGCCT
5551	GGATGCTTTG	AGAGAATTCT	TCAGCACAAT	TGTGGTGGAT	GCCATTGATG
5601	TGTTGAAGTC	CAGGTTTACA	AAGCTAAATG	AATCTACCTT	TGATACTCAA
5651	ATCACCAAGA	AGATGGGCTA	CTATAAGATT	CTAGACGTGA	TGTATTCTCG
5701	CCTTCCCAAA	GATGATGTTC	ATGCTAAGGA	ATCAAAAATT	AATCAAGTTT
5751	TCCATGGCTC	GTGTATTACA	GAAGGAAATG	AACTTACAAA	GACATTGATT
5801	AAATTGTGCT	ACGATGCATT	TACAGAGAAC	ATGGCAGGAG	AGAATCAGCT
5851	GCTGGAGAGG	AGAAGACTTT	ACCATTGTGC	AGCATACAAC	TGCGCCATAT
5901	CTGTCATCTG	CIGIGICTIC	AATGAGTTAA	AATTTTACCA	AGGTTTTCTG
5951	TITAGTGAAA	AACCAGAAAA	GAACTIGCTT	ATTTTTGAAA	ATCTGATCGA
6001	CCTGAAGCGC	CGCTATAATT	TTCCTGTAGA	AGTTGAGGTT	CCTATGGAAA
6051	GAAAGAAAAA	GTACATTGAA	ATTAGGAAAG	AAGCCAGAGA	AGCAGCAAAT
6101	GGGGATTCAG	ATGGTCCTTC	CTATATGTCT	TCCCTGTCAT	ATTTGGCAGA
6151	CAGTACCCTG	AGTGAGGAAA	TGAGTCAATT	TGATTTCTCA	ACCGGAGTTC
6201	AGAGCTATTC	ATACAGCTCC	CAAGACCCTA	GACCTGCCAC	TGGTCGTTTT
6251	CGGAGACGGG	AGCAGCGGGA	CCCCACGGTG	CATGATGATG	TGCTGGAGCT
6301	GGAGATGGAC	GAGCTCAATC	GGCATGAGTG	CATGGCGCCC	CIGACGGCCC
6351	TGGTCAAGCA	CATGCACAGA	AGCCTGGGCC	CGCCTCAAGG	AGAAGAGGAT
6401				AAATTOCTOC	
6451	GGGAAATCCA	ATAGTACCAT	TAAATATCCG	TCTCTTCTTA	GCCAAGCTTG
6501				ACGCGAAGCA	
6551				AATGGAGGAG	
6601				TCTTTCATGG	
6651				TGTTAGCAAA	
6701	AATTICCIAA	TGAAACATGT	CTTTCATCCA	AAAAGAGCTG	TGITTAGACA

Figure 9b (vi)

6751	CAACCTTGAA	ATTATAAAGA	CCCTTGTCGA	GTGCTGGAAG	GATTGTTTAT
6801	CCATCCCTTA	TAGGTTAATA	TTTGAAAAGT	TTTCCGGTAA	AGATCCTAAT
6851	TCTAAAGACA	ACTCAGTAGG	GATTCAATTG	CTAGGCATCG	TGATGGCCAA
6901	TGACCTGCCT	CCCTATGACC	CACAGTGTGG	CATCCAGAGT	AGCGAATACT
6951	TCCAGGCTTT	GGTGAATAAT	ATGTCCTTTG	TAAGATATAA	AGAAGTGTAT
7001	GCCGCTGCAG	CAGAAGTICT	AGGACTTATA	CTTCGATATG	TTATGGAGAG
7051	AAAAAACATA	CTGGAGGAGT	CTCTGTGTGA	ACTGGTTGCG	AAACAATTGA
7101	AGCAACATCA	GAATACTATG	GAGGACAAGT	TTATTGTGTG	CTTGAACAAA
7151	GTGACCAAGA	GCTTCCCTCC	TCTTGCAGAC	AGGTTCATGA	ATGCTGTGTT
7201	CTTTCTGCTG	CCAAAATTTC	ATGGAGTGTT	GAAAACACTC	TGTCTGGAGG
7251	TGGTACTTTG	TCGTGTGGAG	GGAATGACAG	AGCTGTACTT	CCAGTTAAAG
7301	AGCAAGGACT	TCGTTCAAGT	CATGAGACAT	AGAGATGAAA	GACAAAAAGT
7351	ATCTTTCGAC	ATAATTTATA	AGATGATGCC	Aaagttaaaa	CCAGTAGAAC
7401	TCCGAGAACT	TCTGAACCCC	GTTGTGGAAT	TCGTTTCCCA	TCCTTCTACA
7451	ACATGTAGGG	AACAAATGTA	TAATATTCTC	ATGTGGATTC	ATGATAATTA
7501	CAGAGATCCA	GAAAGTGAGA	CAGATAATGA	CTCCCAGGAA	ATATTTAAGT
7551	TGGCAAAAGA	TGTGCTGATT	CAAGGATTGA	TCGATGAGAA	CCCTGGACTT
7601	CAATTAATTA	TTCGAAATTT	CTGGAGCCAT	GAAACTAGGT	TACCTTCAAA
7651	TACCTTGGAC	CCCTTCCTCC	CACTAAATTC	CITATATTCT	CCTAAGATAG
7701	AAGTGCACTT	TTTAAGTTTA	GCAACAAATT	TICIGCTCGA	AATGACCAGC
7751			CCCCATGTTC		
7801			TTGATTCTGA		
7851			GAGACCCAGG		
7901			CCTCTCAGCT		
7951			AGCATGACTT		·
8001			TGGCTGACCG		
8051			ATCTGACTCC		
8101			CACCCTTGAA		
8151			CCAGGGGACG		•
8201			CCTACTACGA		
	GGACCAGGAG				
	AAAAACGAGA				
8351	CAGGTOGTTC	TGTACAGAAG	CTACCGGCAC	GGAGACCTTC	CTGACATTCA

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Figure 9b (vii)

8401	GATCAAGCAC	AGCAGCCTCA	TCACCCCGTT	ACAGGCCGTG	GCCCAGAGGG
8451	ACCCAATAAT	TGCAAAACAG	CTCTTTAGCA	GCTTGTTTTC	TGGAATTTTG
8501	AAAGAGATGG	ATAAATTTAA	GACACTGTCT	GAAAAAAACA	ACATCACTCA
8551	AAAGTTGCTT	CAAGACTTCA	ATCGTTTTCT	TAATACCACC	TTCTCTTTCT
8601	TTCCACCCTT	TGTCTCTTGT	ATTCAGGACA	TTAGCTGTCA	GCACGCAGCC
8651	CTGCTGAGCC	TCGACCCAGC	GGCTGTTAGC	GCTGGTTGCC	TGGCCAGCCT
8701	ACAGCAGCCC	GTGGGCATCC	GCCTGCTAGA	GGAGGCTCTG	CTCCGCCTGC
8751	TGCCTGCTGA	GCTGCCTGCC	AAGCGAGTCC	GTGGGAAGGC	CCGCCTCCCT
8801	CCTGATGTCC	TCAGATGGGT	GGAGCTTGCT	AAGCTGTATA	GATCAATTGG
8851	AGAATACGAC	GLCCLCCGLC	GGATTTTTAC	CAGTGAGATA	GGAACAAAGC
8901	AAATCACTCA	GAGTGCATTA	TTAGCAGAAG	CCAGAAGTGA	TTATTCTGAA
8951	GCTGCTAAGC	AGTATGATGA	GGCTCTCAAT	AAACAAGACT	GGGTAGATGG
9001	TGAGCCCACA	GAAGCCGAGA	AGGATTTTTG	GGAACTTGCA	TCCCTTGACT
9051	GTTACAACCA	CCTTGCTGAG	TGGAAATCAC	TTGAATACTG	TICTACAGCC
9101	AGTATAGACA	GTGAGAACCC	CCCAGACCTA	AATAAAATCT	GGAGTGAACC
9151	ATTTTATCAG	GAAACATATC	TACCTTACAT	GATCCGCAGC	AAGCTGAAGC
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9301	AGAGCTGAGT	CTGCTTTACC	TCCTGCAAGA	TGATGTTGAC	AGAGCCAAAT
. 9351	ATTACATTCA	AAATGGCATT	CAGAGTTTTA	TGCAGAATTA	TICTAGTATT
9401	GATGTCCTCT	TACACCAAAG	TAGACTCACC	AAATTGCAGT	CTGTACAGGC
9451	TTTAACAGAA	ATTCAGGAGT	TCATCAGCTT	TATAAGCAAA	CAAGGCAATT
9501	TATCATCTCA	AGTTCCCCTT	AAGAGACTTC	TGAACACCTG	GACAAACAGA
9551	TATCCAGATG	CTAAAATGGA	CCCAATGAAC	ATCTGGGATG	ACATCATCAC
9601	AAATCGATGT	TTCTTTCTCA	GCAAAATAGA	GGAGAAGCTT	ACCCCTCTTC
9651			_		CAGTGACAGG
9701		AAGAGCAGGA			
9751		ATGAAAATGA			
9801					AGAGTCAAAA
9851					GCCGCCTGAG
9901					CTCACTGTGC
9951					CTACTTAARC
10001			•		GTACAACTTA
10051	CAGGATCATA	GCGAATGCTC	TCAGCAGTGA	GCCAGCCTGC	CITGCTGAAA

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Figure 9b (viii)

1010	1 TCGAGGAGGA CAAGGCTAGA AGAATCTTAG AGCTTTCTGG ATCCAGTTCA
10151	L GAGGATTCAG AGAAGGTGAT CGCGGGTCTG TACCAGAGAG CATTCCAGCA
10201	CCTCTCTGAG GCTGTGCAGG CGGCTGAGGA GGAGGCCCAG CCTCCCTCCT
10251	GGAGCTGTGG GCCTGCAGCT GGGGTGATTG ATGCTTACAT GACGCTGGCA
10301	GATTTCTGTG ACCAACAGCT GCGCAAGGAG GAAGAGAATG CATCAGTTAC
10351	TGATTCTGCA GAACTGCAGG CGTATCCAGC ACTTGTGGTG GAGAAAATGT
10401	TGAAAGCTTT AAAATTAAAT TCCAATGAAG CCAGATTGAA GTTTCCTAGA
10451	TTACTTCAGA TTATAGAACG GTATCCAGAG GAGACTTTGA GCCTCATGAC
10501	AAAAGAGATC TCTTCCGTTC CCTGCTGGCA GTTCATCAGC TGGATCAGCC
10551	ACATGGTGGC CTTACTGGAC AAAGACCAAG CCGTTGCTGT TCAGCACTCT
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10651	CATAAGCAGC GAAAGCTATT CCTTCAAGGA TACTTCTACT GGTCATAAGA
10701	ATAAGGAGTT TGTGGCAAGG ATTAAAAGTA AGTTGGATCA AGGAGGAGTG
10751	ATTCAAGATT TTATTAATGC CITAGATCAG CTCTCTAATC CTGAACTGCT
10801	CTTTAAGGAT TGGAGCAATG ATGTAAGAGC TGAACTAGCA AAAACCCCTG
10851	TAAATAAAA AAACATTGAA AAAATGTATG AAAGAATGTA TGCAGCCTTG
10901	GGTGACCCAA AGGCTCCAGG CCTGGGGGCC TTTAGAAGGA AGTTTATTCA
10951	GACTITITOGA AAAGAATITG ATAAACATIT TGGGAAAGGA GGTTCTAAAC
11001	TACTGAGAAT GAAGCTCAGT GACTTCAACG ACATTACCAA CATGCTACTT
11051	TTAAAAATGA ACAAAGACTC AAAGCCCCCT GGGAATCTGA AAGAATGTTC
11101	ACCCTGGATG AGCGACTTCA AAGTGGAGTT CCTGAGAAAT GAGCTGGAGA
11151	TTCCCGGTCA GTATGACGGT AGGGGAAAGC CATTGCCAGA GTACCACGTG
11201	CGAATCGCCG GGTTTGATGA GCCGGTGACA GTCATGGCGT CTCTGCGAAG
11251	GCCCAAGGGC ATCATCATCC GTGGCCATGA CGAGAGGGAA CACCCTTTCC
11301 11351	TGGTGAAGGG TGGCGAGGAC CTGCGGCAGG ACCAGCGCGT GGAGCAGCTC
11401	TTCCAGGTCA TGAATGGGAT CCTGGCCCAA GACTCCGCCT GCAGCCAGAG
11451	GGCCCTGCAG CTGAGGACCT ATAGCGTTGT GCCCATGACC TCCAGTGATC
11501	CCAGGGCACC GCCGTGTGAA TATAAAGATT GGCTGACAAA AATGTCAGGA
11551	AAACATGATG TTGGAGCTTA CATGCTAATG TATAAGGGCG CTAATCGTAC
11601	TGAAACAGTC ACGTCTTTTA GAAAACGAGA AAGTAAAGTG CCTGCTGATC
	TCTTAAAGCG GGCCTTCGTG AGGATGAGTA CAAGCCCTGA GGCTTTCCTG
11701	GCGCTCCGCT CCCACTTCGC CAGCTCTCAC GCTCTGATAT GCATCAGCCA CTGGATCCTC GGGATTGGAG ACAGACATCT GAACAACTTT ATGGTGGCCA
· -	ACAGACATOLI GAACAACTIT ATGGTGGCCA

Figure 9b (ix)

11751 TGGAGACTGG CGGCGTGATC GGGATCGACT TTGGGCATGC GTTTGGATCC 11801 GCTACACAGT TTCTGCCAGT CCCTGAGTTG ATGCCTTTTC GGCTAACTCG 11851 CCAGTTTATC AATCTGATGT TACCAATGAA AGAAACGGGC CTTATGTACA 11901 GCATCATGGT ACACGCACTC CGGGCCTTCC GCTCAGACCC TGGCCTGCTC 11951 ACCAACACCA TGGATGTGTT TGTCAAGGAG CCCTCCTTTG ATTGGAAAAA 12001 TTTTGAACAG AAAATGCTGA AAAAAGGAGG GTCATGGATT CAAGAAATAA 12051 ATGTTGCTGA AAAAAATTGG TACCCCCGAC AGAAAATATG TTACGCTAAG 12101 AGAAAGTTAG CAGGTGCCAA TCCAGCAGTC ATTACTTGTG ATGAGCTACT 12151 CCTGGGTCAT GAGAAGGCCC CTGCCTTCAG AGACTATGTG GCTGTGGCAC 12201 GAGGAAGCAA AGATCACAAC ATTCGTGCCC AAGAACCAGA GAGTGGGCTT 12251 TCAGAAGAGA CTCAAGTGAA GTGCCTGATG GACCAGGCAA CAGACCCCAA 12301 CATCCTTGGC AGAACCTGGG AAGGATGGGA GCCCTGGATG TGAGGTCTGT 12351 GGGAGTCTGC AGATAGAAAG CATTACATTG TTTAAAGAAT CTACTATACT 12401 TGGTTGGCAG CATTCCATGA GCTGATTTTC CTGAAACACT AAAGAGAAAT 12451 GTCTTTTGTG CTACAGTTTC GTAGCATGAG TTTAAATCAA GATTATGATG 12501 AGTAAATGTG TATGGGTTAA ATCAAAGATA AGGTTATAGT AACATCAAAG

12551 ATTAGGTGAG GITTATAGAA AGATAGATAT CCAGGCTTAC CAAAGTATTA
12601 AGTCAAGAAT ATAATATGTG ATCAGCTTTC AAAGCATTTA CAAGTGCTGC
12651 AAGTTAGTGA AACAGCTGTC TCCGTAAATG GAGGAAATGT GGGGAAGCCT
12701 TGGAATGCCC TTCTGGTTCT GGCACATTGG AAAGCACACT CAGAAGGCTT

12751 CATCACCAAG ATTITGGGAG AGTAAAGCTA

Fig.10.

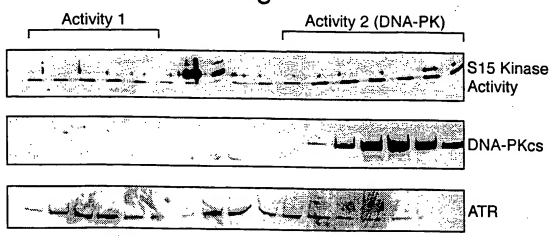
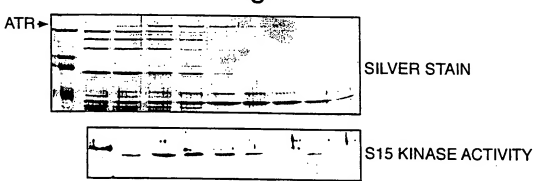


Fig.11.



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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification ⁶: G01N 33/574, 33/68, C12Q 1/48

A3

(11) International Publication Number:

WO 99/04266

(43) International Publication Date:

28 January 1999 (28.01.99)

(21) International Application Number:

PCT/GB98/02115

(22) International Filing Date:

16 July 1998 (16.07.98)

(30) Priority Data:

9714971.0

16 July 1997 (16.07.97)

GB

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(81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, GM, HR, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

Published

With international search report.

Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(88) Date of publication of the international search report:

19 August 1999 (19.08.99)

(54) Title: INTERACTIONS OF ATM, ATR OR DNA-PK WITH P53

(57) Abstract

The interaction of ATM and related protein kinases such as ATR and DNA-PK with p53 is disclosed, in particular the phosphorylation of Ser15 and Thr18 by these proteins. The activity of the proteins is shown to increase in the presence of DNA. Assays for modulators of phosphorylation by the interaction between the proteins and p53 or other proteins having similar phosphorylation sites are provided. Methods of purifying ATM or ATR employing DNA or NTA are also disclosed.

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	actual completion of the international search	"&" document member of	the same patent family
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Name and m	nailing address of the ISA	Authorized officer	
	European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk		
	Tel. (+31-70) 340-2040, Tx. 31 651 epo ni, Fax: (+31-70) 340-3016	Hoekstra	. S
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Inter onal Application No PCT/GB 98/02115

C.(Continu	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	PCT/GB 98/02115
ategory °		
	passages	Relevant to claim No.
X	SAVITSKY K ET AL: "A SINGLE ATAXIA TELANGIECTASIA GENE WITH A PRODUCT SIMILAR TO PI-3 KINASE" SCIENCE,	5,6, 13-16,28
·	vol. 268, 23 June 1996, pages 1749-1753, XP002061971 cited in the application see the whole document	
	see figure 3	
-	SAVITSKY K ET AL: "THE COMPLETE SEQUENCE OF THE CODING REGION OF THE ATM GENE REVEALS SIMILARITY TO CELL CYCLE REGULATORS IN DIFFERENT SPECIES" HUMAN MOLECULAR GENETICS,	5,6, 13-16,28
٠.	vol. 4, no. 11, 1995, pages 2025-2032, XP002061974 cited in the application	
	see the whole document see figure 4	19,20, 24,25
	CIMPRICH K A ET AL: "CDNA CLONING AND GENE MAPPING OF A CONDIDATE HUMAN CELL CYCLE"	29
	PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF USA, vol. 93, April 1996, pages 2850-2855,	
	XP002023632 cited in the application see the whole document see figure 2	
, X	SHIEH, S-Y.: "DNA Damage-induced phosphorylation of p53 alleviates inhibition by MDM2"	1-18
	CELL, vol. 91, 31 October 1997, pages 325-334, XP002095209 see the whole document	
	see page 331, right-hand column, line 27-35	
	WO 98 56391 A (HARVARD COLLEGE ;LEDER PHILIP (US); WESTPHAL CHRISTOPH H (US)) 17 December 1998 see page 12	13-16
	WO 94 12202 A (UNIV DUNDEE ; LANE DAVID PHILIP (GB); HUPP THEODORE ROBERT (GB)) 9 June 1994	1-18
	see the whole document	
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Form PCT/ISA/210 (continuation of second sheet) (July 1992)

Inter .onal Application No PCT/GB 98/02115

C.(Continu	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	PCT/GB 98/02115
Category *		
	:	Relevant to claim No.
A :	HARTLEY K O ET AL: "DNA-DEPENDENT PROTEIN KINASE CATALYTIC SUBUNIT: A RELATIVE OF PHOSPHATIDYLINOSITOL 3-KINASE AND THE ATAXIA TELANGIECTASIA GENE PRODUCT" CELL, vol. 82, 8 September 1995, pages 849-856, XP002065036 cited in the application see the whole document	1-18
A .	ENOCH, T. AND NORBURY, C.: "Cellular responses to DNA damage: cell-cycle checkpoints, apoptosis and the roles of p53 and ATM." TIBS, vol. 20, October 1995, pages 426-430, XP002095210 see the whole document	1-18
Y	A. SUWA, ET AL.: "DNA dependent protein kinase (Ku protein-p350 complex) assembles on double stranded DNA" PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF USA, vol. 91, July 1994, pages 6904-6908, XP002102470 WASHINGTON US see abstract see page 6905, right-hand column, line 20 - page 6907, left-hand column, line 19	19,20, 24,25
A	WO 97 18323 A (ICOS CORP; HOEKSTRA MERL F (US); HOLTZMAN DOUG A (US); KEEGAN KATH) 22 May 1997 see abstract	19,20, 24-27

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Form PCT/ISA/210 (continuation of second sheet) (July 1992)

PCT/GB 98/02115

This International Search Report has not been established in respect of certain claims under Article 17(2(a)) for the following reasons: Claims Nos.:	Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)	
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely: 2. X Claims Nos.: 5.6 (PARTLY) AND 21-23 (ENTIRELY) because they relate to parts of the infamational Application that do not comply with the prescribed requirements to such an extent that no meaningful international Search can be carried out, appecifically: see FURTHER INFORMATION sheet PCT/ISA/210 3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a). Box II Observations where unity of Invention is lacking (Continuation of item 2 of first sheet) This International Searching Authority found multiple inventions in this international application, as follows: see additional sheet 1. X As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee. 3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.: The additional search fees were accompanied by the applicant's protest. Remark on Protest The additional search fees were accompanied by the applicant's protest.	,	
because they relate to subject matter not required to be searched by this Authority, namely: 2.	This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:	
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Form PCT/ISA/210 (continuation of first sheet (1)) (Juty 1998)

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Claims Nos.: 5,6 (PARTLY) AND 21-23 (ENTIRELY)

The subject-matter of claims 5 and 6 in their broadest scope is not defined by technical features that allow the formulation of a meaningful search. These claims do not meet the requirements of Article 6 and Rule 6.3(a) PCT to the extend that a meaningful search is not possible in the entire scope claimed. The subject of the search for hte agents of claims 5 and 6 has been limited to the subject-matter of claims 7-16. As it lies not within the ISA's competence to test a compounds for their capabilites of modulating interactions the scope of the search was formulated by the structural features only. The subject-matter of claims 21-23 is not defined by technical features that allow the formulation of a meaningful search. These claims do not meet the requirements of Article 6 and Rule 6.3(a) PCT to the extend that a meaningful search is not possible

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. Claims: 1-18,28,29

Methods for identifying compounds able to modulate interaction between p53 and DNA binding kinases having the PI3 kinase domain, in particular by phosphorylation of the N-terminus of p53 (ser15) by said kinases, agents so identifiable and medical use of such agents.

2. Claims: 19-27

Methods for identifying compounds able to affect DNA binding by DNA binding kinases having the PI3 kinase domain, agents so identifiable, medical use of such agents and methods involving such binding for purifying ATM or ATR.

information on patent family members

Inte. .onal Application No PCT/GB 98/02115

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